

REPORT DOCUMENTATION PAGE

Form Approved
OMB No. 0704-0188

Public reporting burden for this collection of information is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing this collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to Department of Defense, Washington Headquarters Services, Directorate for Information Operations and Reports (0704-0188), 1215 Jefferson Davis Highway, Suite 1204, Arlington, VA 22202-4302. Respondents should be aware that notwithstanding any other provision of law, no person shall be subject to any penalty for failing to comply with a collection of information if it does not display a currently valid OMB control number. PLEASE DO NOT RETURN YOUR FORM TO THE ABOVE ADDRESS.

1. REPORT DATE (DD-MM-YYYY)	2. REPORT TYPE	3. DATES COVERED (From - To)		
	Technical Paper			
4. TITLE AND SUBTITLE		5a. CONTRACT NUMBER		
<i>Please see attached</i>				
6. AUTHOR(S)		5b. GRANT NUMBER		
7. PERFORMING ORGANIZATION NAME(S) AND ADDRESS(ES)		5c. PROGRAM ELEMENT NUMBER		
<i>ER C</i>				
9. SPONSORING / MONITORING AGENCY NAME(S) AND ADDRESS(ES)		5d. PROJECT NUMBER <i>2303</i>		
Air Force Research Laboratory (AFMC) AFRL/PRS 5 Pollux Drive Edwards AFB CA 93524-7048		5e. TASK NUMBER <i>M2C8</i>		
12. DISTRIBUTION / AVAILABILITY STATEMENT		5f. WORK UNIT NUMBER <i>345709</i>		
Approved for public release; distribution unlimited.		8. PERFORMING ORGANIZATION REPORT		
13. SUPPLEMENTARY NOTES		10. SPONSOR/MONITOR'S ACRONYM(S)		
14. ABSTRACT		11. SPONSOR/MONITOR'S NUMBER(S) <i>Please see attached</i>		
20030129 231				
15. SUBJECT TERMS				
16. SECURITY CLASSIFICATION OF:		17. LIMITATION OF ABSTRACT	18. NUMBER OF PAGES	19a. NAME OF RESPONSIBLE PERSON Leilani Richardson
a. REPORT	b. ABSTRACT	c. THIS PAGE	<i>A</i>	19b. TELEPHONE NUMBER (include area code) (661) 275-5015
Unclassified	Unclassified	Unclassified		

2303M2 CP

MEMORANDUM FOR PRS (In-House Publication)

FROM: PROI (STINFO)

06 March 2002

SUBJECT: Authorization for Release of Technical Information, Control Number: AFRL-PR-ED-TP-2002-048
Karl Christe et al. (PRSP), "Synthesis and Characterization of the SO_2N_3^- , $(\text{SO}_2)_2\text{N}_3^-$ and SO_3N_3^- Anions"

Journal: (2002) Inorganic Chemistry
(Deadline: N/A)

(Statement A)

1. This request has been reviewed by the Foreign Disclosure Office for: a.) appropriateness of distribution statement, b.) military/national critical technology, c.) export controls or distribution restrictions, d.) appropriateness for release to a foreign nation, and e.) technical sensitivity and/or economic sensitivity.

Comments: _____

Signature _____

Date _____

2. This request has been reviewed by the Public Affairs Office for: a.) appropriateness for public release and/or b) possible higher headquarters review.

Comments: _____

Signature _____

Date _____

3. This request has been reviewed by the STINFO for: a.) changes if approved as amended, b) appropriateness of references, if applicable; and c.) format and completion of meeting clearance form if required

Comments: _____

Signature _____

Date _____

4. This request has been reviewed by PR for: a.) technical accuracy, b.) appropriateness for audience, c.) appropriateness of distribution statement, d.) technical sensitivity and economic sensitivity, e.) military/national critical technology, and f.) data rights and patentability

Comments: _____

APPROVED/APPROVED AS AMENDED/DISAPPROVED

PHILIP A. KESSEL
Technical Advisor
Space and Missile Propulsion Division

Date

Synthesis and Characterization of the SO_2N_3^- , $(\text{SO}_2)_2\text{N}_3^-$, and SO_3N_3^- Anions[¶]

Karl O. Christe,^{*†‡} Jerry A. Boatz,[†] Michael Gerken,[‡] Ralf Haiges,[‡] Stefan Schneider,[‡] Thorsten Schroer,[‡] Fook S. Tham,[§] Ashwani Vij,[†] Vandana Vij,[†] Ross I. Wagner,[‡] and William W. Wilson[†]

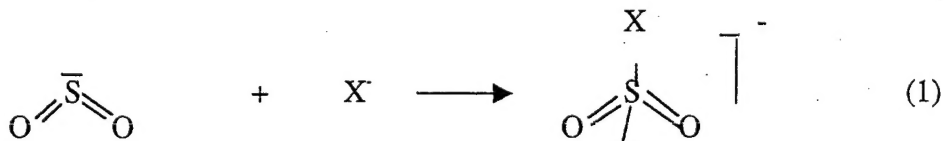
Contribution from the Propulsion Sciences and Advanced Concepts Division, Air Force Research Laboratory (AFRL/PRS), Edwards AFB, California 93524, Loker Hydrocarbon Research Institute and Department of Chemistry, University of Southern California, Los Angeles, California 90089, University of California, Riverside, California 92521

Abstract: SO_2 solutions of azide anions are bright yellow and their Raman spectra indicate the presence of covalently bound azide. Removal of the solvent at -64°C from CsN_3 or $\text{N}(\text{CH}_3)_4\text{N}_3^-$ solutions produces yellow $(\text{SO}_2)_2\text{N}_3^-$ salts. Above -64°C , these salts lose one mole of SO_2 , resulting in white SO_2N_3^- salts that are marginally stable at room temperature and thermally decompose to the corresponding azides and SO_2 . These anions were characterized by vibrational and ^{14}N NMR spectroscopy, and theoretical calculations. Slow loss of the solvent by diffusion through the walls of a sealed Teflon tube containing a sample of $\text{CsSO}_2\text{N}_3^-$ in SO_2 resulted in white and yellowish single crystals that were identified by x-ray diffraction as $\text{CsSO}_2\text{N}_3 \cdot \text{CsSO}_3\text{N}_3$ and $\text{Cs}_2\text{S}_2\text{O}_5 \cdot \text{Cs}_2\text{S}_2\text{O}_7 \cdot \text{SO}_2$, respectively. Pure CsSO_3N_3 was also prepared and characterized by vibrational spectroscopy. The S-N bond in SO_2N_3^- is much weaker than that in SO_3N_3^- , resulting in decreased thermal stability, an increase in the S-N bond distance by 0.23 Å, and an increased tendency to undergo rotational disorder. This marked difference is due to SO_3 being a much stronger Lewis acid (pF value of 7.83) than SO_2 (pF value of 3.99), thus forming a stronger S-N bond with the Lewis base N_3^- . The geometry of the free gaseous SO_2N_3^- anion was calculated at

the RHF, MP2, B3LYP, and CCSD(T) levels. The results show that only the correlated methods correctly reproduce the experimentally observed orientation of the SO_2 group.

Introduction

Sulfur dioxide is a very interesting molecule. It serves not only as a very useful inorganic solvent, but also exhibits a rich reaction-chemistry. In addition to being a ligand with at least 9 different bonding modes,¹ it is also amphoteric and can act either as an electron donor (Lewis base) or acceptor (Lewis acid) molecule. As a donor, it can coordinate through its free valence electron pairs on either sulfur or oxygen. A typical example for the latter case is the oxygen-bridged adduct between SbF_5^- and SO_2 .² As an acceptor, SO_2 likes to expand its coordination number around sulfur from three to four. Typical examples for compounds in which SO_2 acts as an acceptor are the SO_2X^- anions, where X^- stands for a halide or pseudo-halide ion.³



During reactions, aimed at the metathetical synthesis of N_3^+N_3^- , the yellow color of the solutions of colorless N_3^- in colorless SO_2 raised our curiosity. This is in marked contrast to solutions of N_3^- in other polar solvents, such as water, that are colorless. Therefore, the SO_2/N_3^- system was studied in more detail. The results from this study were presented at the 13th European Symposium on Fluorine Chemistry⁴ and a full account of this work is given in this paper. Our original report⁴ provided the first evidence for the existence of the azidosulfite anion and was recently confirmed by an independent crystal structure determination of $[\text{N}(\text{CH}_3)_4]^+[\text{SO}_2\text{N}_3]^-$.⁵ In spite of many known sulfur-nitrogen compounds,¹ including sulfuryl diazide⁶ and azidosulfates,⁷⁻¹² no other reports on azidosulfites were found in the literature.

The SO_3N_3^- anion has previously been prepared by (a) the direct reaction between SO_3 and NaN_3 ,⁸ (b) the reaction of $\text{H}_2\text{NNHSO}_3\text{H}$ and KNO_2 ,⁷ (c) the reaction of $[\text{Na}][\text{SO}_3\text{Cl}]$ and N_3^- ,^{9,12} and (d) the reaction of $\text{S}_2\text{O}_6\text{N}_3^-$ with KOH/NaOH .¹⁰ Despite this diversity of synthetic routes, the SO_3N_3^- anion had been characterized only by elemental analyses and infrared spectroscopy.

Experimental Section

Caution! Azides are highly endothermic and often can decompose explosively. They should be handled on a small scale with appropriate safety precautions.

Materials and Apparatus. Reactions were carried out in either flamed-out Pyrex glass tubes or Teflon-FEP or -PFA ampoules that contained Teflon coated magnetic stirring bars and were closed by stainless steel or Teflon valves. Volatile materials were handled either on a stainless steel / Teflon-FEP vacuum line¹³ or a Pyrex glass vacuum line equipped with grease-free Kontes glass-Teflon valves. Nonvolatile solids were handled in the dry nitrogen atmosphere of a glove box. Infrared spectra were recorded on either a Mattson Galaxy or Midac M Series FT-IR spectrometer using dry powders pressed between AgCl windows in an Econo press (Barnes Engineering Co.). Raman spectra were recorded on either a Bruker Equinox 55 FT-RA spectrometer using a Nd-Yag laser at 1064 nm and Pyrex melting point capillaries, glass NMR or 9 mm Teflon-FEP tubes as sample containers or a Cary Model 83 spectrometer using the 488 nm exciting line of an Ar ion laser. NMR spectra were recorded on a Bruker Avance 400 FT-NMR spectrometer.

The $\text{N}(\text{CH}_3)_4\text{N}_3^-$,¹⁴ CsN_3 ,¹⁵ and CsSO_3Cl ¹⁶ starting materials were prepared by literature methods. The SO_2 (Air Products, anhydrous grade, 99.9%) was dried over CaH_2 . Trimethylsilyl azide (Aldrich, 95%) was condensed from a storage vessel at -35 °C into the reaction vessel at -196 °C.

Preparation of $M^+SO_2N_3^-$ [M = Cs or $N(CH_3)_4$]. In a typical experiment, a 5 mm Pyrex glass NMR tube, closed by a concentric, grease-free Teflon valve (Wilmad Glass Co.) was loaded with CsN_3 (1.335 mmol) in the glove box. The tube was attached to the Pyrex vacuum line and evacuated; then anhydrous SO_2 (0.765 mL) was condensed into the tube at $-196^\circ C$. The mixture was warmed to room temperature, and the CsN_3 dissolved giving a yellow solution. In the presence of traces of moisture, a small amount of a white precipitate formed that was identified by its infrared and Raman spectra as $Cs_2S_2O_5$.¹⁷ The tube and its contents were cooled to $-22^\circ C$, and the SO_2 solvent was pumped off in a dynamic vacuum, leaving behind a white solid (322.0 mg, weight calculated for 1.335 mmol of $CsSO_2N_3$ = 319.1 mg).

Preparation of $M^+(SO_2)_2N_3^-$ [M = Cs or $N(CH_3)_4$]. In a manner similar to that used for the preparation of $CsSO_2N_3$, CsN_3 (1.346 mmol) was loaded into a Pyrex NMR tube equipped with a glass-Teflon valve, and anhydrous SO_2 (0.769 mL) was added in vacuo to the tube at $-196^\circ C$. After warming to room temperature to dissolve all the CsN_3 , the tube was cooled to $-64^\circ C$, and the SO_2 was removed slowly in a dynamic vacuum for 4 hours. The nonvolatile residue consisted of a yellow solid (406 mg, weight calculated for 1.346 mmol of $Cs(SO_2)_2N_3$ = 407.9 mg).

Preparation of $[Cs][SO_3N_3]$. Inside a dry box, a glass vessel, equipped with a Kontes glass-Teflon valve and a Teflon-coated stirring bar, was loaded with $[Cs][SO_3Cl]$ (2.383 mmol). On a glass vacuum line, $(CH_3)_3SiN_3$ (8.940 mmol) was distilled onto the solid at $-196^\circ C$ and allowed to warm to room temperature. After stirring the suspension for two days, volatiles were removed at room temperature yielding a white solid. Raman spectroscopic characterization of the solid revealed the presence of $[Cs][SO_3Cl]$ and $[Cs][SO_3N_3]$. Four additional cycles of adding fresh $(CH_3)_3SiN_3$, followed by removal of all volatiles at room temperature while monitoring the

progress of the reaction by Raman spectroscopy, resulted in a quantitative yield of pure $[\text{Cs}][\text{SO}_3\text{N}_3]$.

Crystal Structure Determination of $\text{CsSO}_2\text{N}_3\cdot\text{CsSO}_3\text{N}_3$. About 0.5 mL of SO_2 was condensed onto anhydrous CsN_3 in a 4 mm FEP NMR tube that was then evacuated at $-196\text{ }^\circ\text{C}$ and heat-sealed. The yellowish solution was allowed to stand for several weeks during which time SO_2 diffused slowly through the walls of the tube, and a mixture of diffraction-quality, prismatic, clear, colorless (compound 1, $\text{CsSO}_2\text{N}_3\cdot\text{CsSO}_3\text{N}_3$) and pale yellow (compound 2, $\text{Cs}_2\text{S}_2\text{O}_5\cdot\text{Cs}_2\text{S}_2\text{O}_7\cdot\text{SO}_2$) crystals was formed. The FEP tube was then cut open under a stream of cold N_2 gas at $\sim -80\text{ }^\circ\text{C}$, and the crystalline contents were dropped into the lip of a low-temperature crystal-mounting apparatus. A Nylon Cryoloop, attached to a magnetic base, was used to trap a crystal, using PFPE (perfluoropolyether) oil, and to mount it on the magnetic goniometer. The single crystal diffraction data were collected on a Bruker 3-circle platform diffractometer, equipped with a SMART¹⁸ CCD (charge coupled device) detector with the χ -axis fixed at 54.74° , and using MoK_α radiation ($\lambda = 0.71073\text{ \AA}$) from a fine-focus tube. This diffractometer was equipped with an LT-3 apparatus for low-temperature data collection using controlled liquid nitrogen boil off. Cell constants were determined from 90 thirty-second frames at $-100\text{ }^\circ\text{C}$ (1) or at $-70\text{ }^\circ\text{C}$ (2). A complete hemisphere of data was collected, using 1271 frames at 30 s/frame at a detector resolution of 512×512 pixels, including 50 frames that were collected at the beginning and end of the data collection to monitor crystal decay. The frames were then processed on a PC running on Windows NT software by using the SAINT software¹⁹ to give the hkl file corrected for Lp/decay. The absorption correction was performed using the SADABS²⁰ program. The structures were solved by the direct method, using the SHELX-97 program,²¹ and refined by the least squares method on F^2 , SHELXL-97,²² incorporated in SHELXTL Suite 5.10

for Windows NT.²³ All atoms were refined anisotropically. For the anisotropic displacement parameters, the U(eq) is defined as one third of the trace of the orthogonalized U_{ij} tensor.

Theoretical Calculations. Theoretical calculations were carried out on IBM RS/6000 work stations using the GAMESS,²⁴ Gaussian 98,²⁵ and ACES II²⁶ program systems and the restricted Hartree Fock (RHF),²⁷ the density functional B3LYP,²⁸ and the correlated MP2²⁹ and single- and double-excitation coupled cluster methods,³⁰ including a non-iterative treatment of connected triple excitations.³¹

Results and Discussion

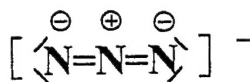
The SO_2/N_3 System. In common solvents, such as water, the N_3^- anion dissolves without color and retains its centro-symmetric, linear $D_{\infty h}$ structure, as shown by its spectroscopic properties. For example, an aqueous NaN_3 solution exhibits in the Raman spectrum only the symmetric N_3^- stretching mode at 1340 cm^{-1} and in the ^{14}N NMR spectrum two resonances at -282.2 and -133.5 ppm, respectively, with an area ratio of 2:1. By contrast, a solution of CsN_3 in liquid SO_2 is intense yellow, exhibits a strong Raman band at 2016 cm^{-1} , and its ^{14}N resonance for the two terminal nitrogen atoms is deshielded by 80 ppm relative to that of the free ion (Table 1). The ^{14}N NMR spectrum of a 1:1 mixture of CsN_3 and SO_2 in CH_3NO_2 solution exhibits also two resonances at $\delta = -209.9$ and -132.5 ppm, respectively, with an area ratio of 2:1. These chemical shifts are similar to those of -201.7 and -133.9 ppm observed for the SO_2 solution. The observation of a single resonance for the terminal nitrogen atoms, even for solutions of the 1:1 and 1:2 adducts in CH_3CN at low temperatures, and the weak temperature dependence of the chemical shifts indicate that the SO_2 groups in azidosulfites undergo rapid exchange on the NMR time scale.

These observations are strong indications that SO_2 forms adducts with the azide anion. The combining ratios of N_3^- with SO_2 were studied both experimentally and by theoretical calculations. It was found that at -64°C SO_2 forms with Cs^+N_3^- yellow 2:1 adducts that lose at about -30°C one mole of SO_2 and yield white 1:1 adducts. The 1:1 adducts are marginally stable at room temperature and, when heated to about 60°C , decompose to pure N_3^- and SO_2 . The corresponding $\text{N}(\text{CH}_3)_4^+$ salts are thermally somewhat less stable. The underlying chemistry can be used for the conversion of insoluble halides, such as fluoride, into azide. For example, CsF is insoluble in most organic solvents and reacts only incompletely with reagents such as trimethylsilylazide. It reacts, however, with SO_2 forming soluble SO_2F . The fluoride in SO_2F can then readily be exchanged for azide using trimethylsilylazide, resulting in the formation of SO_2N_3^- . The latter is then thermally decomposed to give pure azide and SO_2 .¹⁵ During the pyrolysis of SO_2N_3^- , no evidence was obtained for any N_2 elimination before the loss of all the SO_2 , which might have offered a potential synthesis for NSO_2^- salts.³²⁻³⁴ In the presence of small amounts of moisture, colorless cesium salts of $\text{S}_2\text{O}_5^{2-}$ and $\text{S}_2\text{O}_7^{2-}$ can be formed as insoluble by-products.³⁵



As SO_2 has very little vapor pressure below -64°C , the isolation of well defined adducts containing more than two SO_2 molecules per N_3^- could not be studied experimentally.

In agreement with the experimental observations, the theoretical calculations show that SO_2 can form stable 1:1 and 2:1 adducts with N_3^- . In the 1:1 adduct, the SO_2 molecule is attached to one of the terminal nitrogens of the azide anion. This is in accord with the negative charges in N_3^- residing mainly on the terminal nitrogen atoms, thus making them the better donors.



The N-S bond of the resulting adducts is relatively long, varies strongly with the computational level, and ranges for the 1:1 adduct from 1.91 Å at the RHF level to 2.36 Å at the MP2 level.

For the 2:1 adducts, the two SO₂ groups could reside on either the same or both terminal nitrogen atoms. The energy differences between the 1,1 adduct (rN-S = 2.47 Å at MP2) and the 1,3 adduct (rN-S = 2.56 Å) are predicted to be small, i.e., less than 1 kcal/mol at the MP2 and less than 4 kcal/mol at the B3LYP level of theory in favor of the 1,1 adduct.

Based on our calculations, the addition of a third SO₂ molecule to the 1,1 adduct lengthens the two original 1,1 N-S bonds to 2.52 Å, but the third N-S bond to the other terminal nitrogen is much weaker at 2.87 Å. Similarly, the addition of a fourth SO₂ molecule affects the three original N-S bonds relatively little; the two geminal N-S bonds at 2.54 Å become slightly longer and the third N-S bond at 2.80 Å becomes slightly shorter; however, the fourth N-S bond at 3.19 Å becomes much longer. These calculations at the MP2 level confirm that in the presence of excess SO₂, the N-S bonds become very weak, thus accounting for the observed facile N-S bond breakage on an NMR timescale. A more detailed discussion of the calculated structures will be given below.

The SO₂N₃⁻ anion can be described as either a donor-acceptor adduct between the Lewis acid SO₂ and the Lewis base N₃⁻ or, by analogy to many known SO₂X⁻ (X = halogen or pseudohalogen) anions, as an azidosulfite anion. Whereas either description is acceptable, the description of the higher (SO₂)_nN₃⁻ adducts as azidopolysulfites is inappropriate because there is no significant interaction between the SO₂ ligands. For the analogous (SO₃)_nN₃⁻ adducts, azidopolysulfate structures have previously been suggested;¹⁰ however it appears likely that these SO₃ analogues also contain discrete SO₃ units and not polysulfate anions.

Crystal structures of CsSO_2N_3 and CsSO_3N_3 . The white crystals (compound 1), obtained from a sample of CsSO_2N_3 in SO_2 solution that was kept in a Teflon NMR tube at room temperature for several weeks, had the composition $\text{CsSON}_3 \cdot \text{CsSO}_3\text{N}_3$. These crystals were stable only at low-temperature or under an atmosphere of SO_2 at room temperature. The asymmetric unit in the crystal lattice of 1 shows two Cs, two sulfur and five oxygen atoms besides six nitrogen atoms forming two azide groups (Figure 1 and Tables 2-4). The cesium and sulfur atoms, and O1 and N2 lie on special positions. The large thermal ellipsoids observed for the N1, N3, N4, N5 and N6 atoms indicate a disorder across the crystallographic mirror plane. These atoms were then moved off the mirror plane and refined which resulted in two disordered azide conformations on the sulfur groups. In SO_2N_3^- , the sulfur atom S2 is connected to three oxygen positions with a total occupancy of 2 and, therefore, is attributed to an SO_2 group. The oxygen atoms on S2 exhibit rotational disorder and have refined occupancies for O3 and O5 of 64 and 36 %, respectively. A comparison of the geometry found in this study for SO_2N_3^- in its cesium salt (see Figure 2) with that in the ordered $\text{N}(\text{CH}_3)_4^+$ salt⁵ and the theoretical predictions at the correlated level (see Figure 3 and Table 4) shows good agreement and demonstrates that the counter ion does not strongly influence the structure of SO_2N_3^- .

The SO_3N_3^- anion is also slightly disordered. The N1 and N3 atoms are disordered across the crystallographic symmetry plane through S1, O1, and N2 (see Figure 4). Figure 5 depicts one of the two disordered components. Furthermore, the rotation of the SO_3 group with respect to the N_3 group deviates from the predictions at the MP2 and B3LYP levels of theory (see Table 5 and Figure 6). In the free gaseous anion, the O1, S1, N1, N2, and N3 atoms are all predicted to be coplanar, with O1 being *trans* to the azide group, while in the experimental solid-state structure it is *cis*. Since the calculated energy difference between the *cis*- and *trans*-rotamers is very small

due to the weak S-N bond and its low frequency (33 cm^{-1}) torsion mode, the orientation of the SO_3 group is easily affected by the strong oxygen-cesium interactions in the lattice (see below).

A comparison of the experimental structures of SO_2N_3^- and SO_3N_3^- shows a remarkable difference of 0.23 \AA for the S-N distances, i. e., $1.981(12)\text{ \AA}$ for SO_2N_3^- and $1.754(10)\text{ \AA}$ and SO_3N_3^- . The observed difference agrees well with that of 0.32 \AA predicted at the B3LYP level and can be rationalized by the large difference in the Lewis acidities of SO_2 and SO_3 . Based on the Christe/Dixon pF scale,³⁶ SO_2 and SO_3 have pF values of 3.99 and 7.83, respectively. If one views the SO_2N_3^- and SO_3N_3^- anions as Lewis acid / Lewis base adducts (see above), it is not surprising that the stronger Lewis acid SO_3 forms the more stable adduct with the azide anion. Due to its longer S-N bond, the SO_2N_3^- anion should undergo SO_n group rotation even more easily than SO_3N_3^- . Therefore, one can expect the orientations of the SO_n groups in the SO_nN_3^- anions to vary significantly from one salt to another. The $\text{N}_\beta\text{-N}_\gamma$ distances at $\sim 1.13\text{ \AA}$ are significantly shorter than the $\text{N}_\alpha\text{-N}_\beta$ distances at $\sim 1.21\text{ \AA}$ and are characteristic for covalent azides with significant triple and double bond character, respectively. With increasing S-N bond length and a weakening of the $\text{SO}_2\text{-N}_3^-$ interaction, the $\text{N}_\alpha\text{-N}_\beta$ and $\text{N}_\beta\text{-N}_\gamma$ bonds become, as expected, more similar and their values approach those of the free N_3^- anion.

The $\text{CsSO}_2\text{N}_3\cdot\text{CsSO}_3\text{N}_3$ structure exhibits many short contacts indicative of strong interionic interactions. Thus, all the Cs...O distances fall within the range of $3.047(6)$ ($\text{Cs2}\cdots\text{O1}$) to $3.40(2)$ ($\text{Cs2}\cdots\text{O5}$) \AA and are considerably shorter than the sum of their van der Waals radii of about 4.5 \AA .³⁷ The Cs...N contacts of $3.219(13)$ and $3.305(11)\text{ \AA}$, involving the SO_2N_3^- and the SO_3N_3^- groups, respectively, are also significantly shorter than the sum of their van der Waals radii of about 4.54 \AA . The $\text{S1}\cdots\text{Cs2}$ contact of $3.848(2)\text{ \AA}$ is also considerably shorter than the average distance of $\sim 4.43\text{ \AA}$ for the remaining Cs...S contacts.

The short Cs...O contacts mentioned above result in a three-dimensional network of four-membered Cs_2O_2 rings involving alternating SO_3N_3^- and SO_2N_3^- units (Figure 1). A mean least square plane analysis of these two four-membered rings shows that they are twisted by 93.4° with respect to each other. The cesium-sulfur network (Figure 7) forms the basic crystal lattice that is built up of face- and edge-sharing cuboids stacked in a stair-like fashion.

The yellow impurity **2** was identified as a double salt of $\text{Cs}_2\text{S}_2\text{O}_5$ and $\text{Cs}_2\text{S}_2\text{O}_7$ with a molecule of SO_2 present as a solvate. The crystal structures of the separate anions have previously been reported,³⁸⁻⁴¹ and the structure of this ternary compound will be discussed in a forthcoming paper.

Vibrational Spectra and Theoretical Calculations. The vibrational spectra of $\text{CsSO}_2\text{N}_3^-$, $\text{N}(\text{CH}_3)_4\text{SO}_2\text{N}_3^-$, $\text{Cs}(\text{SO}_2)_2\text{N}_3^-$, $\text{N}(\text{CH}_3)_4(\text{SO}_2)_2\text{N}_3^-$, and $\text{CsSO}_3\text{N}_3^-$ are summarized in Tables 6-8 and Figures 8-11. The assignments were made by comparison with the calculated spectra and are supported by normal coordinated analyses. The correctness of the given mode descriptions was established by the calculated potential energy distributions. All the listed calculations were carried out with a 6-311+G(d) basis set. The calculations at the B3LYP level were also done with 6-311+G(2d) and aug-cc-pvtz basis sets and showed that the choice of the basis set had little influence on the results. For the ease of presentation, the spectra are discussed in the following order: SO_2N_3^- , $(\text{SO}_2)_2\text{N}_3^-$, N_3^- in SO_2 solution, and SO_3N_3^- .

SO_2N_3^- . The structure of the SO_2N_3^- anion is well established by the crystal structures of its $\text{N}(\text{CH}_3)_4^{+5}$ and Cs^+ (Fig. 2) salts and the results from the theoretical calculations (Figure 3 and Table 4). It must be emphasized that only the calculations at a correlated level (Figure 3) duplicate the experimentally observed orientation of the SO_2 group. At the RHF level, the SO_2 group points in the same direction as the azido group (Fig. 12). However, the energy difference

between the structures of Figures 3 and 12 is only about one kcal/mol because of the rotational barrier along the N-S bond in SO_2N_3^- ($\nu = 22 \text{ cm}^{-1}$ at the B3LYP level) being even lower than that in SO_3N_3^- ($\nu = 33 \text{ cm}^{-1}$). The agreement between the observed and the calculated spectra (see Table 6) is satisfactory if one keeps in mind that the skeletal modes involving the very long N-S bonds are strongly correlation dependent. Thus, the uncorrelated RHF method underestimates the bond length of the S-N bond resulting in high frequency values for the modes involving the N-S bond. On the other hand, all the correlated methods significantly overestimate the S-N bond length (see Tables 4 and 5) resulting in low frequency values for these modes. If one assumes that the actual bond lengths and frequencies fall between the uncorrelated and the correlated values, the fit between calculated and observed data becomes satisfactory. These results, together with our previous study of the SO_2F^- anion,⁴² demonstrate the difficulties associated with the accurate calculation of highly ionic, long bonds. In view of the significant discrepancies between the observed and calculated frequencies, the calculated force fields have not been included in this paper.

$(\text{SO}_2)_2\text{N}_3^-$. Whereas the structure of the SO_2N_3^- anion is firmly supported by its crystal structures, only Raman data are available for the $(\text{SO}_2)_2\text{N}_3^-$ ion. Since the negative charges in N_3^- are located on the two terminal nitrogen atoms (see above), the SO_2 groups must be attached to these. However, the two SO_2 groups could be attached to the same or to different nitrogen atoms. Theoretical calculations at the B3LYP (Figures 13 and 15) and MP2 (Figures 14 and 16) levels of theory favor the geminal 1,1- $(\text{SO}_2)_2\text{N}_3^-$ over the terminal 1,3- $\text{O}_2\text{SNNNSO}_2^-$ adduct by 3.5 and 0.4 kcal/mol, respectively. Figures 13 and 14 also show that the B3LYP and MP2 calculations result in very different minimum energy structures for the 1,3-adduct. At the MP2 level there is, in addition to the $\text{N}_\alpha\text{-S}$ bond with 2.56 Å, also significant $\pi\text{-}\pi$ interaction between the N=N and

S=O double bonds resulting in an N_{β} -O interaction of 2.84 Å, while at the B3YLP level the interaction occurs mainly through N-S bonds. Calculations were also carried out using the MP2 structure as the starting point for the B3LYP calculation and *vice versa*, but in each case the calculations reverted back to the structures shown in Figures 13 and 14, establishing that they are the true minima for each method. For the 1,1-adducts, the MP2 and B3LYP structures are more similar (Figures 15 and 16).

Distinction between the 1,1- and the 1,3- structures is possible from the observed Raman intensities (see Table 7). In the 1,3-structures the relative intensities of the antisymmetric and the symmetric azide stretching modes are comparable, while in the 1,1 adducts the antisymmetric stretching mode is much more intense than the symmetric one. Furthermore, in the 1,3 adduct many of the low-frequency skeletal modes are of low Raman intensity, while in the 1,1-adduct they are more intense and of comparable intensity. The observed spectra (see Table 7 and Figure 10) clearly favor the energetically favored 1,1-structure.

SO₂ Solutions of N₃⁻. The Raman spectra of the yellow solutions of N₃⁻ in liquid SO₂ show bands at about 2016(s) ($\nu_{as}N_3^-$), 1274(w) ($\nu_{as}SO_2$), 1120(vs) ($\nu_{sym}SO_2$), 646(mw) (δN_3^-), 400(br), and 230(br) cm⁻¹ that are in accord with the spectra observed for the yellow, solid 1,1-(SO₂)₂N₃⁻ adduct. In view of the similar spectra and the fact that the theoretical calculations indicate that the bonding of a third and fourth SO₂ ligand to (SO₂)₂N₃⁻ becomes increasingly weaker (see above), the yellow species present in the N₃⁻/SO₂ solutions is attributed to the 1,1-(SO₂)₂N₃⁻ anion.

SO₃N₃⁻. The structure of this anion is firmly established by the x-ray diffraction results (Figures 4 and 5). The theoretical calculations at the MP2 and B3LYP levels (see Tables 5 and 8 and Figure 6) result in almost identical geometries, however, compared to the experimental

structure, the SO_3 group is rotated by 60° about the S-N axis. As for SO_2N_3^- , both methods significantly overestimate the N-S bond length, in this case by 0.12 \AA . This bond distance overestimate results in an underestimate of those vibrational frequencies involving the S-N bond (see Table 8), but otherwise the agreement between observed and calculated spectra is satisfactory. The fact that the S-N bond in SO_3N_3^- is significantly shorter than that in SO_2N_3^- has already been explained above by the increased Lewis acidity of SO_3 . This increased withdrawal of electron density and negative charge from the azide anion in SO_3N_3^- also causes a marked decrease in the N-N bond lengths, an increase in the average N_3^- stretching and bending frequencies, and an increased bond length difference between the $\text{N}_\alpha\text{-N}_\beta$ and $\text{N}_\beta\text{-N}_\gamma$ bonds.

Conclusions. The N_3^- anion acts as a pseudohalide and dissolves in liquid SO_2 forming yellow donor-acceptor adducts containing at least two SO_2 molecules per N_3^- . At low temperature, yellow $(\text{SO}_2)_2\text{N}_3^-$ salts can be isolated in which both SO_2 molecules are probably attached to the same terminal nitrogen atom. At about -30°C , one mole of SO_2 can be pumped off, resulting in marginally stable, colorless SO_2N_3^- salts that can be converted at about 50°C into pure azides. The azidosulfites are analogous to the previously known azidosulfates, but possess a significantly weaker S-N bond resulting in lowered thermal stability.

Acknowledgements

The authors thank the National Science Foundation, the Defense Advanced Project Agency, and the Air Force Office of Scientific Research for financial support. S.S. thanks the Alexander von Humboldt Foundation for a Feodor-Lynen fellowship and M. G. the Natural Sciences and Engineering Research Council of Canada for a postdoctoral fellowship. This work was also supported in part by a grant of IBM SP time from the Aeronautical Systems Center at

Wright-Patterson Air Force Base, a Department of Defense High Performance Computing Major Shared Resource Center.

Supporting Information Available

Tables of structure determination summary, atomic coordinates, bond lengths and angles and anisotropic displacement parameters of CsSO_2N_3 , CsSO_3N_3 in CIF format. This material is available free of charge via the internet at <http://pubs.acs.org>.

References

[¶]This paper is dedicated to Professor Dieter Naumann on the occasion of his 60th birthday.

[†]Air Force Research Laboratory

[‡]University of Southern California

[§]UC Riverside

- (1) Greenwood, N. N.; Earnshaw, A. *Chemistry of the Elements*, Second Edition, Butterworth-Heinemann, Oxford, 1998, p. 721-746.
- (2) Bacon, J.; Dean, P. A. W.; Gillespie, R. J. *Can. J. Chem.* 1969, 47, 1655.
- (3) Kornath, A.; Blecher, O.; Ludwig, R. *J. Am. Chem. Soc.* 1999, 121, 4019, and references cited therein.

(4) Vij, A.; Wilson, W.; Sheehy, J.; Boatz, J.; Gerken, M.; Schneider, S.; Schroer, T.; Haiges, R.; Wagner, R.; Tham, F.; Christe, K.; Paper B7, presented at the 13th European Symposium on Fluorine Chemistry, Bordeaux, France, July 15-20, 2001.

(5) Kornath, A.; Blecher, O.; Ludwig, R. *Z. Anorg. Allg. Chem.* **2002**, *628*, 183.

(6) Curtius, T.; Schmidt, F. *Ber. Dtsch. Chem. Ges.* **1922**, *55*, 1571.

(7) Traube, W.; Vockerodt, A. *Ber. Dtsch. Chem. Ges.* **1914**, *47*, 943.

(8) Beck, G. *J. Prakt. Chem.* **1940**, *156*, 227.

(9) Elsner, H.; Ratz, H. *German Patent* 886,298 **1953**.

(10) Lehmann, H. A.; Holznagel, W. *Z. Anorg. Allg. Chem.* **1958**, *293*, 314.

(11) Ruff, J. K. *Inorg. Chem.* **1965**, *4*, 567.

(12) Shozda, R. J.; Vernon, J. A. *J. Org. Chem.* **1967**, *32*, 2876.

(13) Christe, K. O.; Wilson, W. W.; Schack, C. J.; Wilson, R. D. *Inorg. Synth.* **1986**, *24*, 39.

(14) Christe, K. O.; Wilson, W. W.; Bau, R.; Bunte, S. W. *J. Am. Chem. Soc.* **1992**, *114*, 3411.

(15) Gerken, M.; Schneider, S.; Schroer, T.; Christe, K. O. *Z. Anorg. Allg. Chem.* in press.

(16) Ciruna, J. A.; Robinson, E. A. *Can. J. Chem.* **1968**, *46*, 1715.

(17) Siebert, H. Anwendungen der Schwingungsspektroskopie in der Anorganischen Chemie, Anorganische und Allgemeine Chemie in Einzeldarstellungen, VII; Springer Verlag: Berlin, 1966, p 102; Herlinger, A. W.; Long, T. V. *Inorg. Chem.* **1969**, *8*, 2661.

(18) SMART V 4.045, Software for the CCD Detector System, Bruker AXS, Madison, WI, 1999.

(19) SAINT V 4.035, Software for the CCD Detector System, Bruker AXS, Madison, WI, 1999.

(20) SADABS, Program for absorption correction for area detectors, Version 2.01, Bruker AXS, Madison, WI (2000).

(21) Sheldrick, G. M. SHELXS-97, Program for the Solution of Crystal Structure, University of Göttingen, Germany, 1997.

(22) Sheldrick, G. M. SHELXL-97, Program for the Refinement of Crystal Structure, University of Göttingen, Germany, 1997.

(23) SHELXTL 5.10 for Windows NT, Program library for Structure Solution and Molecular Graphics, Bruker AXS, Madison, WI (1997).

(24) Schmidt, M.W.; Baldridge, K.K.; Boatz, J.A.; Elbert, S.T.; Gordon, M.S.; Jensen, J.H.; Koseki, S.; Matsunaga, N.; Nguyen, K.A.; Su, S.J.; Windus, T.L.; Dupuis, M.; Montgomery, J.A. *J. Comp. Chem.* 1993, 14, 1347-1363.

(25) Frisch, M.J.; Trucks, G.W.; Schlegel, H.B.; Scuseria, G.E.; Robb, M.A.; Cheeseman, J.R.; Zakrzewski, V.G.; Montgomery, J.A., Jr.; Stratmann, R.E.; Burant, J.C.; Dapprich, S.; Millam, J.M.; Daniels, R.E.; Kudin, K.N.; Strain, M.C.; Farkas, O.; Tomasi, J.; Barone, V.; Cossi, M.; Cammi, R.; Mennucci, B.; Pomelli, C.; Adamo, C.; Clifford, S.; Ochterski, J.; Peterson, G.A.; Ayala, P.Y.; Cui, Q.; Morokuma, K.; Malick, D.K.; Rabuck, A.D.; Raghavachari, K.; Foresman, J.B.; Cioslowski, J.; Ortiz, J.V.; Stefanov, B.B.; Liu, G.; Liashenko, A.; Piskorz, P.; Komaromi, I.; Gomperts, R.; Martin, R. L.; Fox, D. J.; Keith, T.; Al-Laham, M. A.; Peng, C.Y.; Nanayakkara, A.; Gonzalez, C.; Challacombe, M.; Gill, P.M.W.; Johnson, B.; Chen, W.; Wong, M.W.; Andres, J.L.; Gonzalez, C.; Head-Gordon, M.; Replogle, E.S.; Pople, J.A. *Gaussian 98*, revision A.6; Gaussian, Inc.: Pittsburgh, PA, 1998.

(26) Stanton, J.F.; Gauss, J.; Watts, J.D.; Nooijen, M.; Oliphant, N.; Perera, S.A.; Szalay, P.G.; Lauderdale, W.J.; Gwaltney, S.R.; Beck, S.; Balkova, A.; Bernholdt, D.E.; Baeck,

K.K.; Rozyczko, P.; Sekino, H.; Hober, C.; Bartlett, R.J. *ACES II, Quantum Theory Project*; University of Florida: Integral packages included are VMOL (Almlöf, J.; Taylor, P.R.), BPROPS (Taylor, P.R.), and ABACUS (Helgaker, T.; Jensen, H.J.Aa.; Jorgensen, P.; Olsen, J.; Taylor, P.R.)

(27) Levine, I. N. *Quantum Chemistry*, 3rd ed.; Allyn and Bacon: Boston, 1983:p 373.

(28) The B3LYP functional uses a three-parameter exchange functional of Becke (B3) [Becke, A.D. *J. Chem. Phys.* **1993**, *98*, 5648; Stephens, P.J.; Devlin, C.F.; Chabalowski, C.F.; Frisch, M.J. *J. Phys. Chem.* **1994**, *98*, 11623] and the Lee, Yang, and Parr (LYP) correlation gradient-corrected functional [Lee, C.; Yang, W.; Parr, R.G. *Phys. Rev. B* **1988**, *37*, 785].

(29) a) Pople, J.A.; Binkley, J.S.; Seeger, R. *Int. Quantum Chem.* **1976**, *10*, 1.
b) Bartlett, R.J.; Silver, D.M. *Int. Quantum Chem.* **1975**, *9*, 183.
c) Dupuis, M.; Chin, S.; Marquez, A. in: G. Malli (ed.), *Relativistic and Electron Correlation Effects in Molecules*, Plenum, New York, 1994.
d) Frisch, M.J.; Head-Gordon, M.; Pople, J.A. *Chem. Phys. Lett.* **1990**, *166*, 275.
e) Bartlett, R.J.; Stanton, R.J. Applications of post-Hartree-Fock methods: A Tutorial, in: Lipkowitz, K.B.; Boyd, D.B. (Eds.), *Reviews of Computational Chemistry*, Vol. V, VCH Publishers, New York, 1994.

(30) Purvis, G.D., III; Bartlett, R.J. *J. Chem. Phys.* **1982**, *76*, 1910.

(31) Raghavachari, K.; Trucks, G.W.; Pople, J.A.; Head-Gordon, M. *Chem. Phys. Lett.* **1989**, *157*, 479.

(32) Roesky, H. W.; Schmieder, W.; Isenberg, W.; Boehler, D.; Sheldrick, G. M. *Angew. Chem., Int. Ed. Engl.* **1982**, *21*, 153; Roesky, H. W.; Panday, K. P.; Krebs, B.; Dartmann, M. *J. Chem. Soc., Dalton Trans.* **1984**, 2271.

(33) Chivers, T.; Schmidt, K. J.; McIntyre, D. D.; Vogel, H. J. *Can. J. Chem.* **1989**, *67*, 1788.

(34) Morgan, N. H.; Linnert, H. V.; Riveros, J. M. *J. Phys. Chem.* **1995**, *99*, 11667.

(35) Seel, F.; Mueller, E. *Chem. Ber.* **1955**, *88*, 1747.

(36) Christe, K. O.; Dixon, D. A.; McLemore, D.; Wilson, W. W.; Sheehy, J. A.; Boatz, J. A. *J. Fluorine Chem.* **2000**, *101*, 151.

(37) Bondi, A. *J. Phys. Chem.* **1964**, *68*, 441.

(38) Lindqvist, I. *Acta Crystallogr.* **1957**, *10*, 406.

(39) Baggio, S. *Acta Crystallogr.* **1971**, *27*, 517.

(40) Wang, Y.; Chen, I-C. *Acta Crystallogr., Sect. C* **1984**, *40*, 1780.

(41) Hvoslef, J.; Tracy, M. L.; Nash, C. P. *Acta Crystallogr., Sect. C* **1986**, *42*, 353.

(42) Lork, E.; Mews, R.; Viets, D.; Watson, P. G.; Borrman, T.; Vij, A.; Boatz, J. A.; Christe, K. O. *Inorg. Chem.* **2001**, *40*, 1303.

Table 1. ^{14}N NMR Spectra of Aqueous and SO_2 Solutions of MN_3 ($\text{M} = \text{Na, Cs or N(CH}_3)_4$) and of MSO_2N_3 ($\text{M} = \text{Cs or N(CH}_3)_4$) in CH_3NO_2 , CH_3CN , or CHF_3 Solution

compound (solvent)	chem shift, ppm (line width, Hz)		
	central	terminal nitrogen	area ratio
NaN_3 (H_2O , 20 °C)	-133.5 (18)	-282.2 (60)	1:2
NaN_3 (SO_2 , 20 °C) ^b	-135.8 weak	not observed	
CsN_3 (SO_2 , 20 °C) ^c	-133.9 (15)	-201.7 (76)	1:2
(SO_2 , -25 °C)	-134.3 (19)	-200.1 (150)	1:2
(SO_2 , -65 °C)	-134.7 (51)	-198.9 (235)	1:2
CsSO_2N_3 (CH_3NO_2 , -22 °C)	-132.5 (16)	-209.9 (155)	1:2
$\text{N}(\text{CH}_3)_4\text{N}_3$ (H_2O , 22 °C) ^d	-133.4 (18)	-281.0 (58)	1:2
$\text{N}(\text{CH}_3)_4\text{N}_3$ (SO_2 , -65 °C) ^d	-135.0 (35)	-199.8 (200)	1:2
$\text{N}(\text{CH}_3)_4\text{SO}_2\text{N}_3$ (CH_3CN , -40 °C) ^{e,f}	-132.7 ^f	-205.3 (140)	
$\text{N}(\text{CH}_3)_4\text{SO}_2\text{N}_3$ (CHF_3 , -90 °C) ^b	-136.4 weak	not observed	
$\text{N}(\text{CH}_3)_4(\text{SO}_2)_2\text{N}_3\text{nSO}_2$ (CH_3CN , -40 °C) ^{c,f}	-133.1 ^f	-203.7 (192)	

^aClear colorless solution with faint yellow solid. ^bLow solubility. ^cPale yellow solution with yellow solid. ^dThe sharp resonances due to $\text{N}(\text{CH}_3)_4^+$ were observed at -139.3 (H_2O) and -336.9 ppm (SO_2). ^eResonances due to CH_3CN and $\text{N}(\text{CH}_3)_4^+$ were observed at -137.7 (~300Hz) and -338 ppm, respectively. ^f Not determined due to overlap with solvent resonance.

Table 2. Crystal Data and Structure Refinement for $\text{CsSO}_2\text{N}_3\text{CsSO}_3\text{N}_3$ (1)

empirical formula	$\text{Cs}_2\text{N}_6\text{O}_5\text{S}_2$
fw	494.00
Temperature (K)	173(2)
space group	$P2_1/m$
a , Å	9.542(2)
b , Å	6.2189(14)
c , Å	10.342(2)
β , deg	114.958(4)
V , Å ³	556.4(2)
Z	2
ρ_{calc} g/cm ³	2.949
μ , mm ⁻¹	6.939
crystal size, mm	0.10 x 0.27 x 0.52
λ , Å	0.71073
R_{int}	0.0446
transmission factors	0.5437- 0.1230
goodness-of-fit on F	1.075
$R1$, $wR2$ [$I > 2\sigma(I)$]	0.0398, 0.1068
$R1$, $wR2$ (all data)	0.0416, 0.1085

Table 3. Atomic Coordinates (x 10⁴) and Equivalent Isotropic Displacement Parameters (Å² x 10³) for CsSO₂N₃CsSO₃N₃. U(eq) is Defined as one Third of the Trace of the Orthogonalized U^{ij} Tensor.

	x	y	z	U(eq)
Cs(1)	6199(1)	2500	3672(1)	38(1)
Cs(2)	10539(1)	2500	1983(1)	39(1)
S(1)	6914(2)	2500	7536(2)	42(1)
S(2)	9094(3)	-2500	3854(2)	40(1)
O(1)	8336(7)	2500	8827(6)	59(2)
O(2)	6625(5)	562(8)	6701(5)	49(1)
O(3)	8278(16)	-3010(20)	4683(12)	47(5)
O(4)	8677(11)	-276(12)	3266(9)	45(2)
O(5)	10543(17)	-3070(30)	3910(30)	45(7)
N(1)	5404(10)	1975(19)	8035(10)	49(4)
N(2)	5694(7)	2500	9245(6)	31(1)
N(3)	5843(12)	3101(19)	10317(11)	51(4)
N(4)	7990(17)	-4208(17)	2097(14)	55(3)
N(5)	7791(13)	-3134(14)	1059(12)	40(2)
N(6)	7645(14)	-2130(30)	100(13)	52(5)

Table 4. Most Important Bond Lengths [Å] and Angles [°] of the SO_2N_3^- Anion in its Cs^+ Salt Compared to Those in $[\text{N}(\text{CH}_3)_4]^+\text{SO}_2\text{N}_3^-$ and the Calculated^a V Values

	calculated			observed		
	RHF	B3LYP	MP2	CCSD(T)	$\text{Cs}^+\text{SO}_2\text{N}_3^-$	$[\text{N}(\text{CH}_3)_4]^+\text{SO}_2\text{N}_3^-$
bond distances (Å)						
S-N	1.907	2.226	2.356	2.198	1.981(12)	2.005(2)
S-O ^c	1.464	1.492	1.492	1.494	1.410(10) ^e	1.453(2)
S-O ^d	1.462	1.490	1.491	1.493	1.495(8)	1.457(2)
N α -N β	1.205	1.207	1.223	1.222	1.209(16)	1.214(2)
N β -N γ	1.113	1.169	1.205	1.182	1.130(16)	1.144(2)
bond angles (°)						
O-S-N ^d	101.46	99.21	99.94	99.01	100.1(5)	96.82(8)
O-S-N ^e	102.09	102.42	102.24	101.46	102.7(11) ^f	100.87(9)
S-N α -N β	112.0	110.29	106.81	108.67	110.0(8)	110.22(14)
N α -N β -N γ	178.9	179.16	179.32	179.37	178.2(15)	178.6(2)
O-S-O	112.6	114.58	115.58	114.90	111.1(9) ^e	112.72(10)

^aA 6-31+G(d) basis set was used for all calculations. ^bData from ref 2. ^cOxygen atom that is approximately perpendicular to the plane of the N_3S fragment. ^dOxygen atom that is approximately in plane with the N_3S fragment. ^eAverage value from disordered positions. ^fTaken from the major component of the disordered oxygen position.

Table 5. Most Important Bond Lengths [Å] and Angles [°] of the SO_3N_3^- Anion in its Cs^+ Salt Compared to the Calculated* Values

	calculated	observed	
	B3LYP /6-31+G(d)	MP2 /6-31+G(d)	
bond distances (Å)			
S-N	1.876	1.871	1.754(10)
S-O'	1.473	1.472	1.440(5)
S-O ₂	1.479	1.478	1.447(6)
N α -N β	1.223	1.233	1.207(11)
N β -N γ	1.151	1.175	1.120(11)
bond angles (°)			
O'-S-N	99.59	99.31	92.7(4)
O ₂ -S-N	102.94	102.47	107.2(4)
S-N α -N β	114.21	114.52	110.0(8)
N α -N β -N γ	175.39	174.48	173.5(10)
O'-S-O ₂	116.18	116.45	114.9(2)
S-N-N-N	180.00	180.00	180.0
O'-S-N-N	180.00	180.00	180.0

Table 6. Vibrational Spectra of Solid $[N(CH_3)_4][SO_2N_3]$ and $C_8[SO_2N_3]$ and Assignment of the $SO_2N_3^-$ Anion Bands Based on the Unscaled Frequencies and Intensities Calculated at the RHF, MP2, B3LYP, and CCSD(T) Levels of Theory.^a

RHF	MP2	B3LYP	CCSD(T)	vibrational frequencies, cm^{-1}			assignments	
				calcd for free $SO_2N_3^-$	exp	C_8^+ IR, 22 °C	Raman, 22 °C	Raman, -100 °C
375(1456)[146]	2143(889)[107]	2106(1182)[120]	2043	3039(45)				
1239(3)[34]	1350(42)[17]	1291		3035(40)sh				
[289(431)[7.3]	1244(249)[56]	1198(297)[25]	1202	2995(11)sh				
164(207)[20]	1087(199)[203]	1065(266)[65]	1056	2981(13)sh				
1143(2)				2958(36)				
1179(2)				2923(11)				
1188(7)				2915(10)				
1196s				2883(4)				
1190(9)				2859(4)				
1272m				2824(2)sh				
1274(13)				2817(5)				
1188(7)				2797(3)				
1197(5)				2019(19)	2036vs	2036 (20)	2042 (16)	(1333+428+269)
1404(2)				2014 (27)	2013 (25)	2018(30)	$\nu_{as}(N_3)$	$\nu_2(A_1), \delta_s(CH_3)$
1332(2)				1488(2)				$\nu_6(E), \delta_{as}(CH_3)$
1417(2)				1469(13)				$\nu_{16}(F_2), \delta_s(CH_3)$
1404(2)				1463(15)				
1329m				1417(2)				
1330(5)				1404(2)				
1333(5)				1332(2)				
(2x663)				1294(2)				
1274(14)				1272(7)				
1188(9)				1197(5)				
1179(2)				1188(7)				
1143(2)				1143(2)				
1083(47)				1073s		1077(100)	1145(1)	$\nu_s(SO_2)$
1076(20)sh							1078(61)	
952(12)								$\nu_{18}(F_2), \nu_{as}(CN_4)$
758(9)								$\nu_3(A_2), \nu_s(CN_4)$

Table 6, continued....

Table 6, continued....

RHF	MP2	B3LYP	CCSD(T)	vibrational frequencies, cm^{-1}			SO ₂ N ₃ ⁻	N(CH ₃) ₄ ⁺	assignments
				calcd for free SO ₂ N ₃ ⁻	exp	Cs ⁺ IR, 22 °C	Raman, 22 °C		
793(10)[10]	586(1)[4]	646(6)[1]	598	665(0+)	662mw	660(3)	663(2)	$\delta(\text{N}_3)$ in SN ₃ plane	
700(18)[0.1]	550(1)[1]	608(6)[3]	549	544(4)	547m	540(7)	543(5)	$\delta(\text{N}_3)$ out of SN ₃ plane	
607(116)[1.4]	497(33)[4]	504(27)[4]	505	460(5)	414ms	420(15)	428(17)	$\delta_{\text{sciss}}(\text{SO}_2)$	$\nu_{19}(\text{F}_2), \delta(\text{CN}_4)$
499(263)[6.4]	329(133)[10]	369(122)[8]	368	405(12)	377(2)	309(30)	317(45)	$\delta_{\text{s}}\text{SO}_2\text{N}$	
352(13)[2.9]	181(11)[12]	220(1)[6]	223	295(10)	224(100)	258(84)	269(100)	$\tau(\text{SO}_2)$	
310(189)[29]	167(38)[17]	181(50)[20]	177	154(28)	169(20)	175(24)	175(24)	$\nu(\text{SN})$	
150(4.0)[11]	69(4)[11]	106(9)[14]	96	100(24)	125sh	96(100)	125sh	$\delta_{\text{sciss}}(\text{SNN})$	
				86(17)sh	103(56)	86sh	103(56)	lattice vibrations	
				78(31)		78(27)			
17(0.9)[8.4]	30(1)[10]	22(1)[7]	25					$\tau(\text{O}_2\text{S}-\text{N}_3)$	

*Bands arising from the FEP sample tube were observed at 735 (2) cm^{-1} in the spectrum of the N(CH₃)₄⁺ salt.

Table 7. Vibrational Spectra of $[N(CH_3)_4][SO_2]_2N_3$ and $[Cs][SO_2]_2N_3$ and Assignments for $(SO_2)_2N_3$ Based on the Frequencies and Intensities Calculated at the MP2 and B3LYP Levels of Theory.^{a,b}

vibrational frequencies, cm^{-1}		assignments				
$O_2SNNNSO_2$	calcd for free	$(O_2S)_2\text{NNN}$		$(O_2S)_2N_3N(CH_3)_4^+$		
MP2	B3LYP	B3LYP	MP2	exp		
2096(10)[93]	2076(35)[11]	2125(21)[137]	2102(12)[206]	3052(8)sh 3043(12) 3037(8)sh 2992(13) 2976(5) 2960(6) 2930(10) 2903(1) 2826(4)	$N(CH_3)_4^+C_3^+$ Raman, -80 °C ^c Raman, -100 °C	$\nu(CH_3)$ modes
1241(10)[46]	1379(0)[22]	1339(2)[11]	1252(4)[58]	1981(13) 1492(1) 1471(1) 1455(6) 1422(1) 1383(1) 1333(13)	2028(1) 2028(27) 2019(17)	$\nu_2(A_1)$, $\delta_1(CH_3)$ $\nu_4(E)$, $\delta_4(CH_3)$
1236(1)[67]	1243(11)[29]	1235(5)[24]	1242(5)[38]	1250(1)	$\nu_1(F_2)$, $\delta_1(CH_3)$	
1228(8)[13]	1241(0)[17]	1224(5)[24]	1217(0,1)[32]	1246(2) 1223(3)		
1052(1)[457]	1094(4)[307]	1088(2)[176]	1080(4)[287]	1177(1) 1140(100)	$\nu_6(N_3)$	
1052(5)[143]	1077(7)[112]	1078(6)[51]	1067(2)[219]	1136(98) 1128(30) 1082(1)	$\nu_7(E)$, CH_3 rock	
				1120(100) 1113(6) 1099(41) 1096sh	$\nu_4(SO_2)$	

Table 7. continued...

O ₂ SNNNNSO ₂		vibrational frequencies, cm ⁻¹				assignments
		calcd for free (O ₂ S) ₂ NNN		N(CH ₃) ₄ ⁺ CS ⁺ Raman, -80 °C ^a		
MP2	B3LYP	B3LYP	MP2	exp	(O ₂ S) ₂ N(CH ₃) ₄ ⁺	
563(0)[2]	640(0)[4]	638(0)[2]	563(0.1)[0.6]	1074(1)	v ₁₈ (F ₂), v ₁₈ (CN ₄)	
546(0)[1]	620(0)[1]	606(0)[2]	502(0.3)[1.4]	956(4)	v ₃ (A ₂), v ₄ (CN ₄)	
488(1)[0]	501(1)[1]	504(1)[2]	491(0.1)[5]	756(6)		
487(0)[5]	498(0)[0]	503(0)[5]	487(1)[5]	644(2)		
				617(1.6)		
				546(0.4)		
				535(5)	δ _{as} (SO ₂)	
				458(2)	v ₁₈ (F ₂), δ(CN ₄)	
				384(1)	v ₁₂ (F ₁), τ(CH ₃)	
				370(1)		
				352(2)		
276(2)[8]	306(0)[1.5]	323(2)[6]	312(2)[5]	292(1)		
261(1)[5]	281(5)[1]	308(4)[4]	276(3)[1.5]	231(7)		
201(0)[10]	209(0)[6]	247(1)[3]	221(1)[9]	254(4)		
201(0)[15]	172(0)[3]	181(0)[8]	162(0.4)[2]	206(5)		
140(0)[5]	131(0)[10]	146(1)[4]	139(0.4)[7]	183(4)		
135(1)[0]	130(2)[1]	112(0)[1.5]	109(0.1)[7]	161(14)		
109(0)[1]	71(0)[4]	78(0)[4]	79(0.03)[9]	128(10)		
97(0)[2]	52(0)[3]	73(0)[6]	68(0.08)[6]	117(5)		
71(0)[5]	20(0)[10]	34(0)[2]	45(0.01)[3]	105(9)		
47(0)[1]	19(0)[1]	30(0)[3]	26(0.05)[2]	105(11)		
				89(5)		
				78(5)		
				69(4)		
34(0)[0]	1(0)[1]	22(0)[0]	23(0.03)[2]	54(4)		

^aBands arising from the FEP sample tube were observed at 734 (3) cm⁻¹ for the N(CH₃)₄(O₂S)₂N₃ sample. ^bRaman spectra of Cs(O₂S)₂N₃ recorded at -130 °C on the Cary instead of the Bruker Equinox showed significant changes in the relative intensities of some of the bands.

Table 8. Vibrational Spectra of $[\text{Cs}][\text{SO}_3\text{N}_3]$

vibrational frequencies, cm^{-1}				assignments
exp		calc		SO_3N_3 (C_s)
IR, 22 °C	Raman, 22 °C	MP2	B3LYP	
3337m				2134+1247 ($\nu_1 + \nu_{11}$)
2469m				1269+1247 ($\nu_3 + \nu_{11}$)
2450m				2x1247 (2x ν_{11})
2283w				1247+1045 ($\nu_4 + \nu_{11}$)
2134vs	2138(23) 2118(15)	2272(19)[81]	2192(827)[89]	$\nu_1(\text{A}')$, $\nu_{as}(\text{N}_3)$
1483w	1278(8)	1285(1)[1]	1323(136)[8]	2x744 (2x ν_5)
1269vs		1283(9)[16]	1241(318)[14]	$\nu_2(\text{A}')$, $\nu_s(\text{N}_3)$
1247vs	1250(9)	1261(9)[7]	1218(336)[9]	$\nu_3(\text{A}')$, $\nu_{as}(\text{SO}_3)$
1114w	1089(5)			$\nu_{11}(\text{A}'')$, $\nu_{as}(\text{SO}_3)$
1045vs	1052(100)	1022(4)[44]	987(158)[40]	$\nu_4(\text{A}')$, $\nu_s(\text{SO}_3)$
1028w				562+464 ($\nu_{13} + \nu_9$)
744s	746(18)	706(0.5)[5]	700(18)[5]	$\nu_s(\text{A}')$, $\delta(\text{N}_3)$ in plane
640sh	644(10)	558(0.1)[0.1]	593(8)[0.1]	$\nu_{12}(\text{A}'')$, $\delta(\text{N}_3)$ out of plane
plane		563(9)[7]	572(349)[6]	$\nu_6(\text{A}')$, $\delta(\text{SO}_3)$
626vs				
umbrella				2x313 (2x ν_4)
618m				
579w				
562s	558(13)	519(1)[4]	510(28)[4]	$\nu_{13}(\text{A}'')$, $\delta_{as}(\text{SO}_3)$
556s				
547s	550sh	518(0.5)[3]	509(18)[3]	$\nu_7(\text{A}')$, $\delta_{sciss}(\text{SO}_2)$
464m	464(54)	363(2)[29]	369(34)[24]	$\nu_9(\text{A}')$, $\nu(\text{S}-\text{N})$
	370(28)	314(1)[11]	314(11)[9]	$\nu_8(\text{A}')$, $\delta_{rock}(\text{SO}_3)$
364sh				
313(8)		322(0)[1]	318(0)[1]	$\nu_{14}(\text{A}'')$, $\delta_{wag}(\text{SO}_3)$
185(23)		123(0)[7]	139(1)[6]	$\nu_{10}(\text{A}')$, $\delta_{sciss}(\text{S}-\text{N}-\text{N})$
112sh		49(0)[6]	33(0)[6]	$\nu_{15}(\text{A}'')$, $\tau(\text{S}-\text{N})$

Diagram Captions

Figure 1. Crystal Structure of $\text{CsSO}_2\text{N}_3\text{CsSO}_3\text{N}_3$.

Figure 2. A 30% Ortep plot of the experimental SO_2N_3^- structure.

Figure 3. Structure of the SO_2N_3^- anion calculated at the CCSD(T)/6-31+G(d) level.

Figure 4. Crystal structure of the SO_3N_3^- anion in its Cs salt, showing the disorder of N1 and N3 across the crystallographic mirror plane.

Figure 5. A 30% Ortep plot of the experimental SO_3N_3^- structure.

Figure 6. Structure of the SO_3N_3^- anion calculated at the MP2/6-31+G(d) level.

Figure 7. Packing diagram of $\text{CsSO}_2\text{N}_3\text{CsSO}_3\text{N}_3$ showing the cesium-sulfur network.

Figure 8. Raman spectrum of CsSO_2N_3 .

Figure 9. Raman spectrum of $\text{N}(\text{CH}_3)_4\text{SO}_2\text{N}_3$.

Figure 10. Raman spectrum of $\text{Cs}(\text{SO}_2)_2\text{N}_3$.

Figure 11. Infrared (upper trace) and Raman (lower trace) spectra of CsSO_3N_3 .

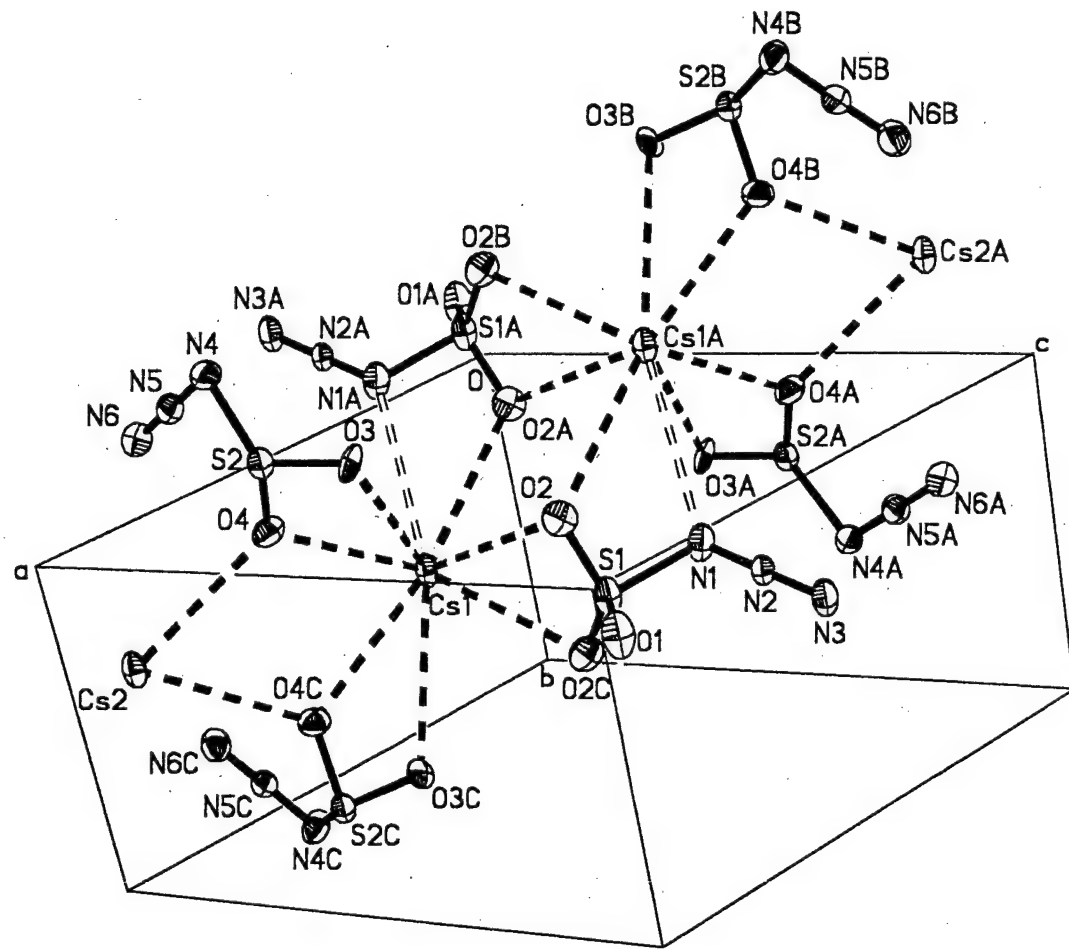
Figure 12. Structure of the SO_2N_3^- anion calculated at the RHF/6-31+G(d) level.

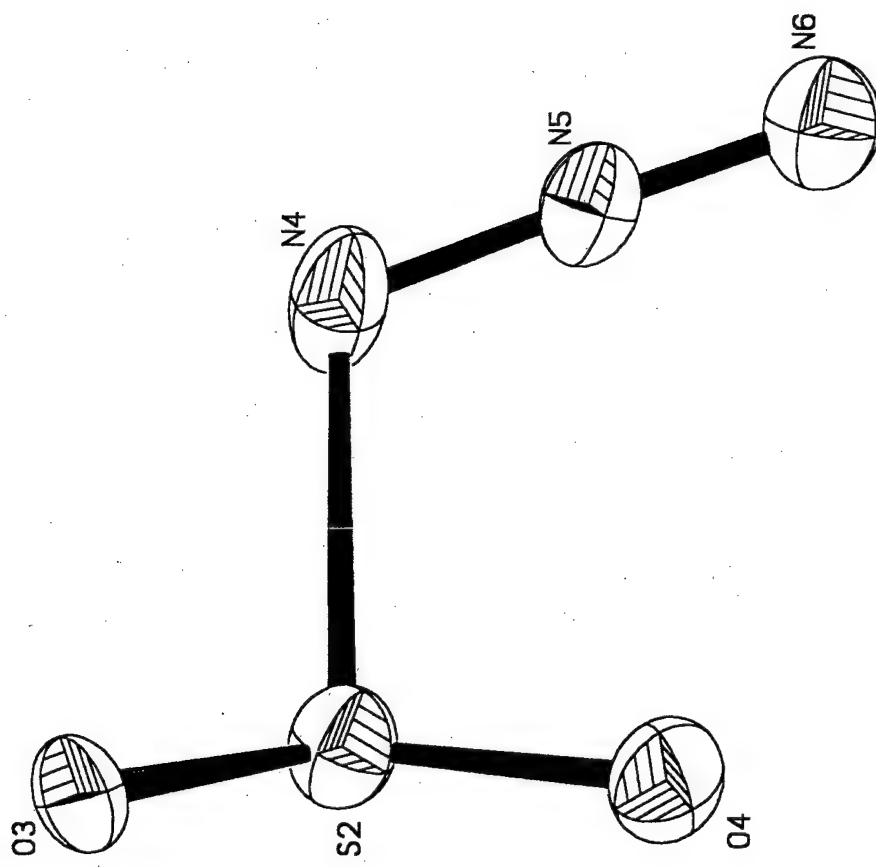
Figure 13. Structure of the 1,3-(SO_2)₂ N_3^- anion calculated at the B3LYP/6-31+G(d) level.

Figure 14. Structure of the 1,3-(SO_2)₂ N_3^- anion calculated at the MP2/6-31+G(d) level.

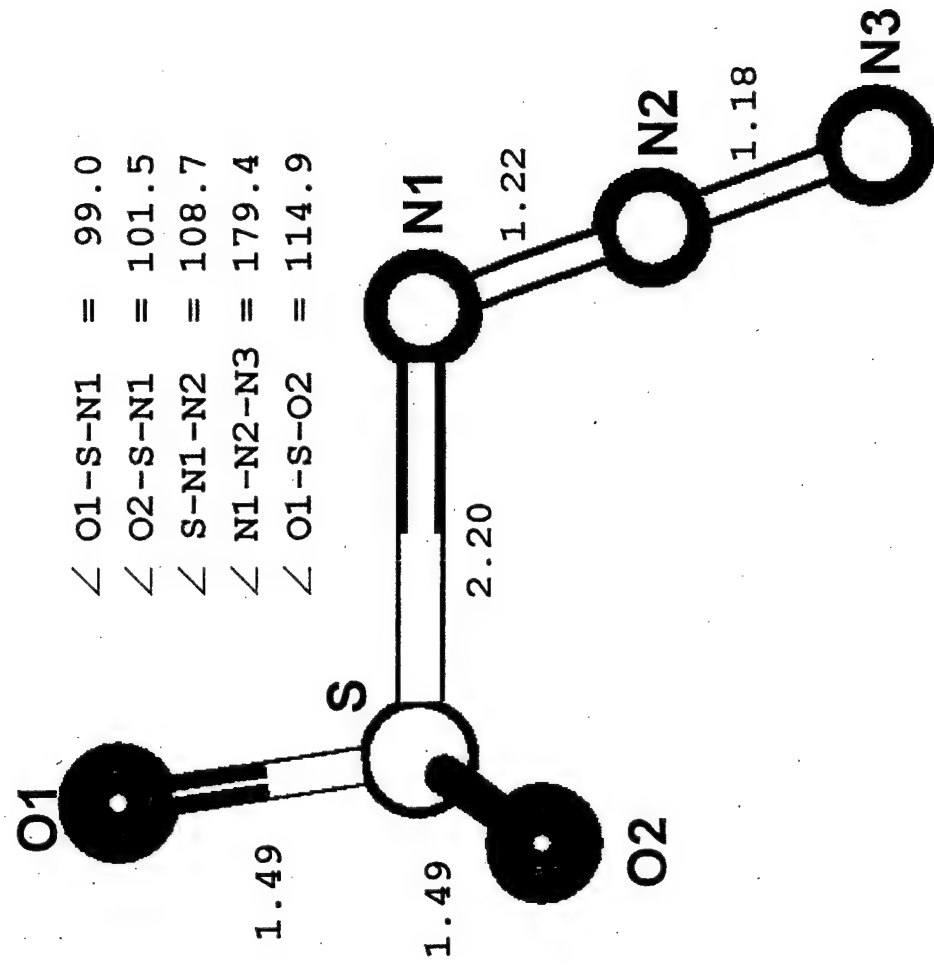
Figure 15. Structure of the 1,1-(SO_2)₂ N_3^- anion calculated at the B3LYP/6-31+G(d) level.

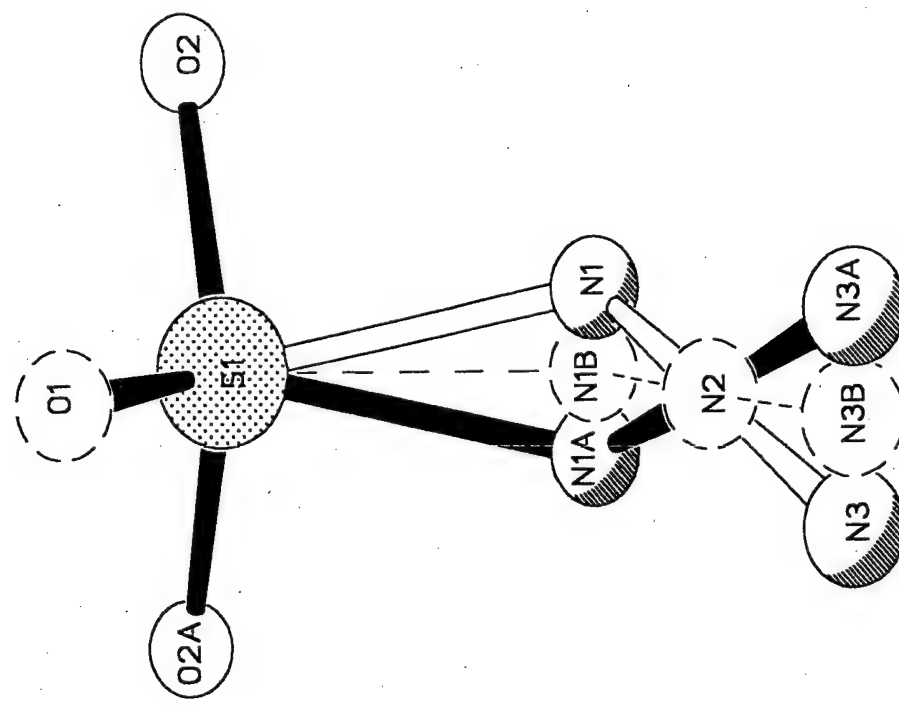
Figure 16. Structure of the 1,1-(SO_2)₂ N_3^- anion calculated at the MP2/6-31+G(d) level.

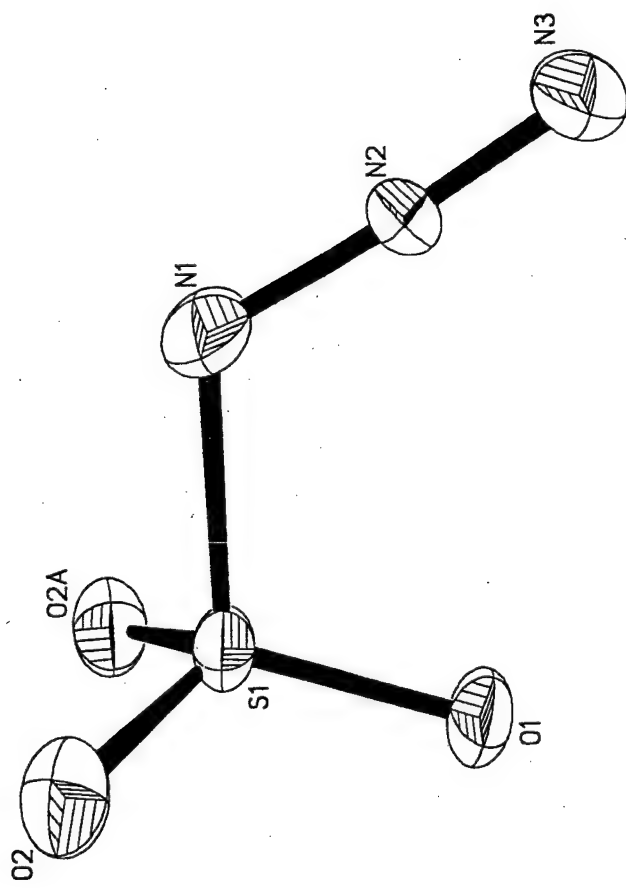




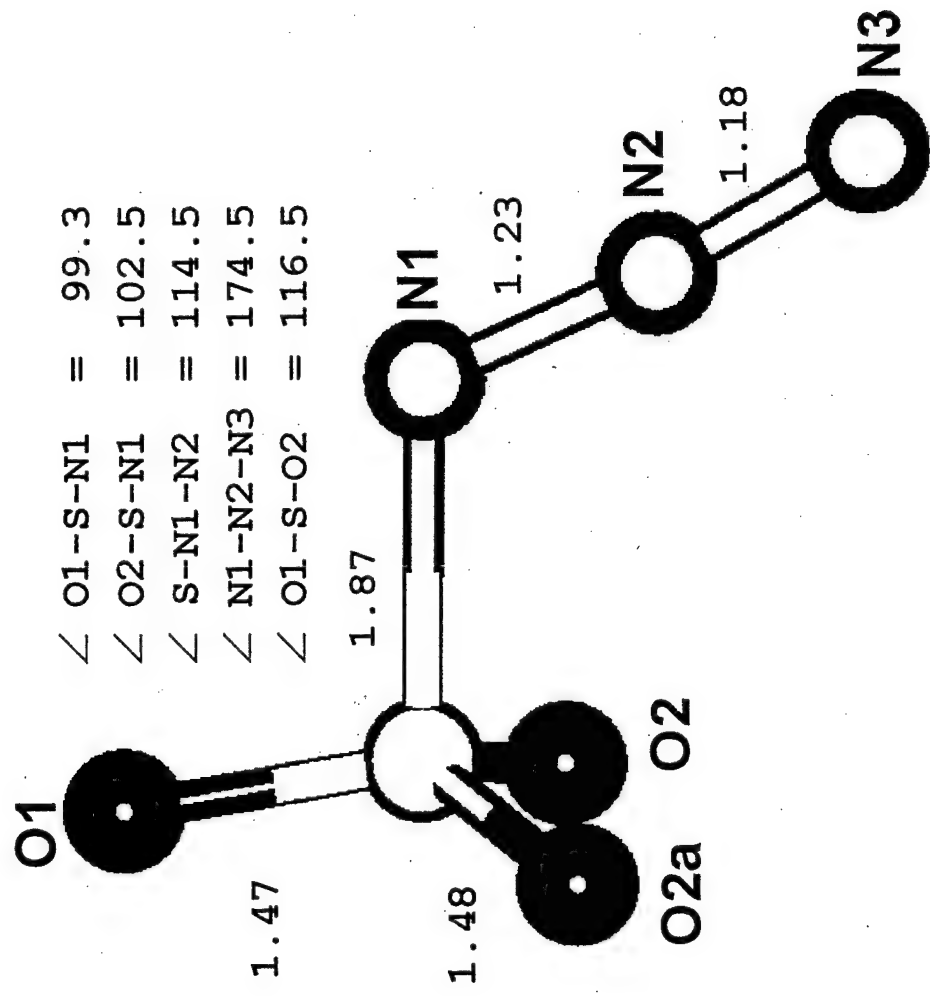
CCSD(T) / 6-31+G(d)

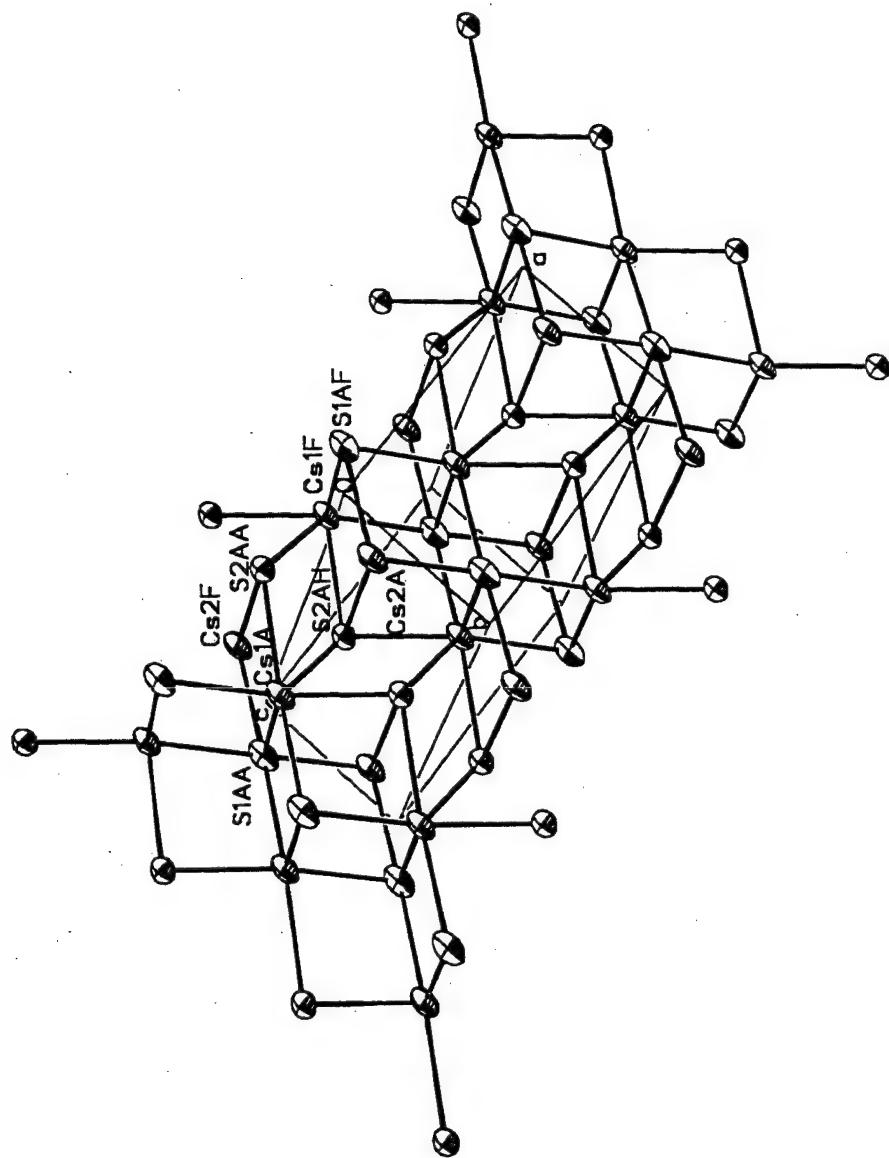


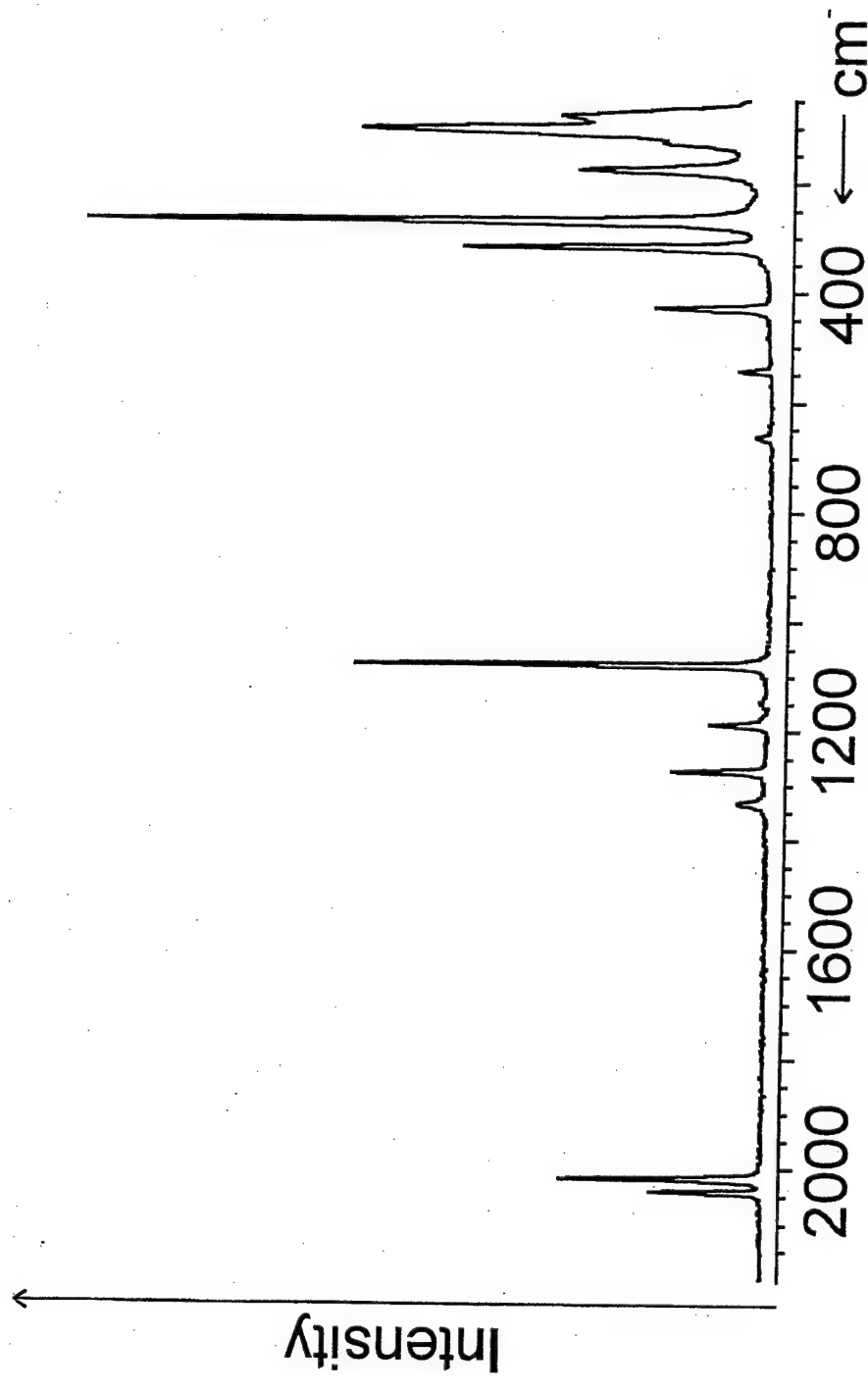


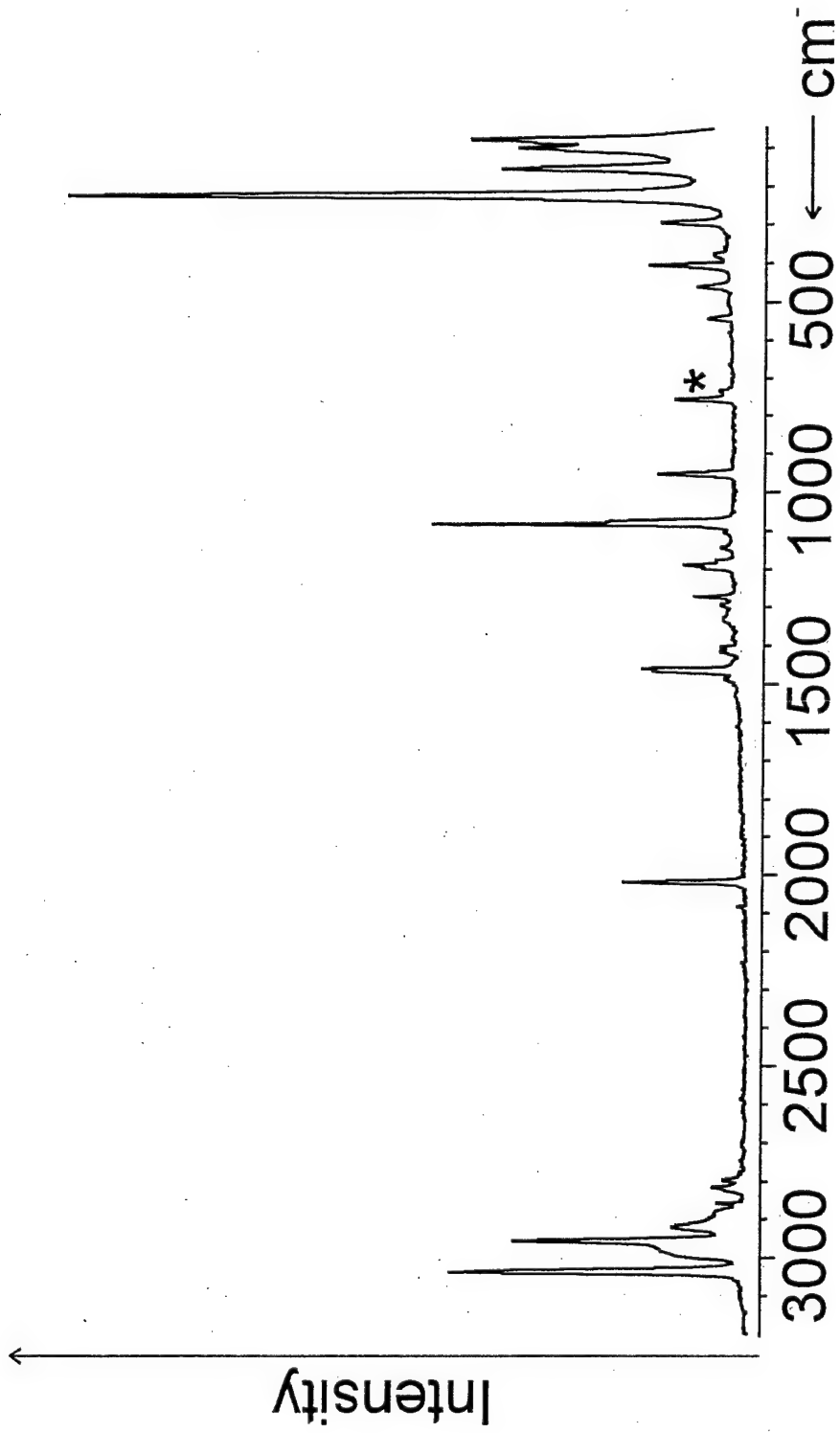


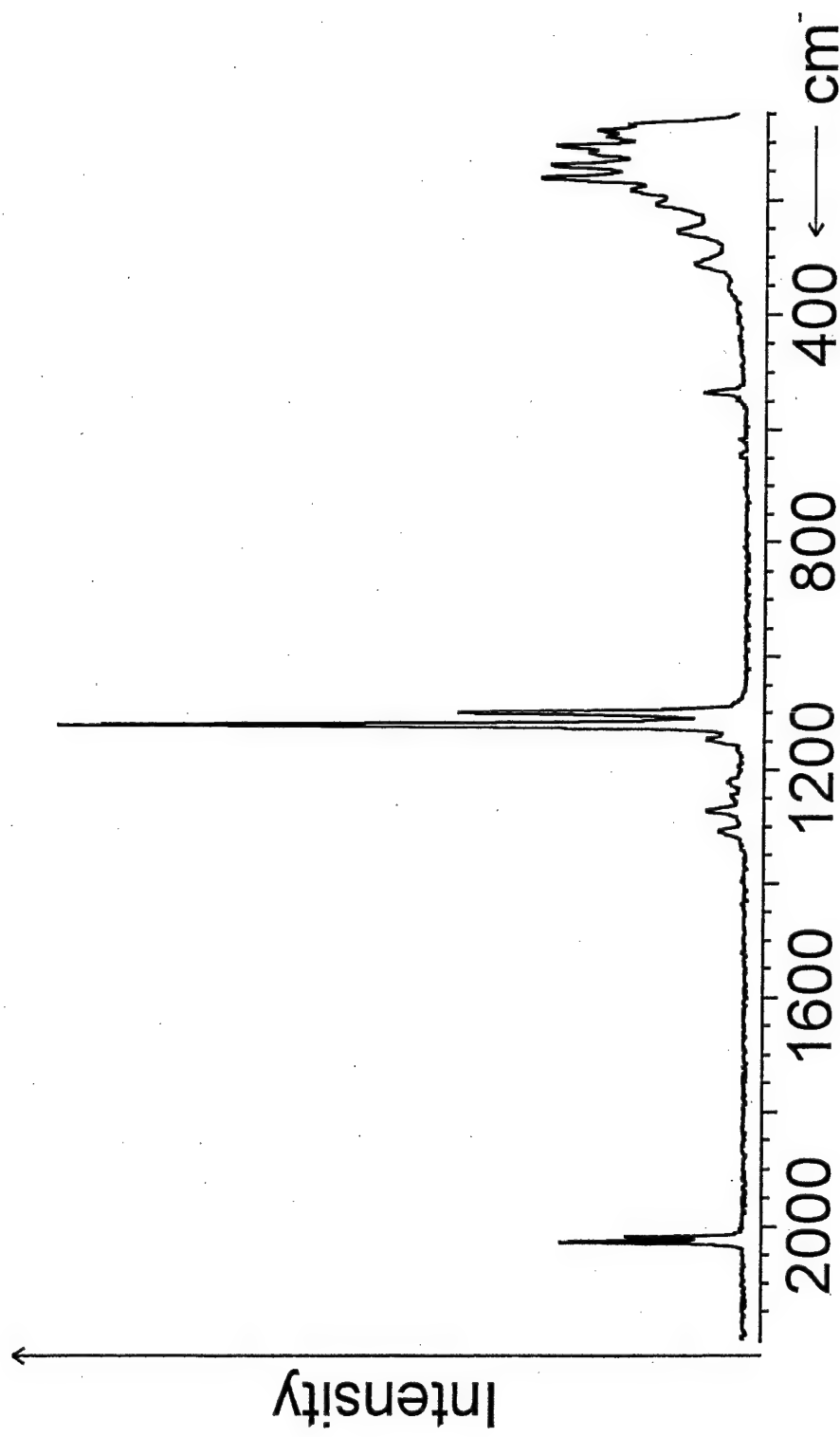
MP2/6-31+G(d)

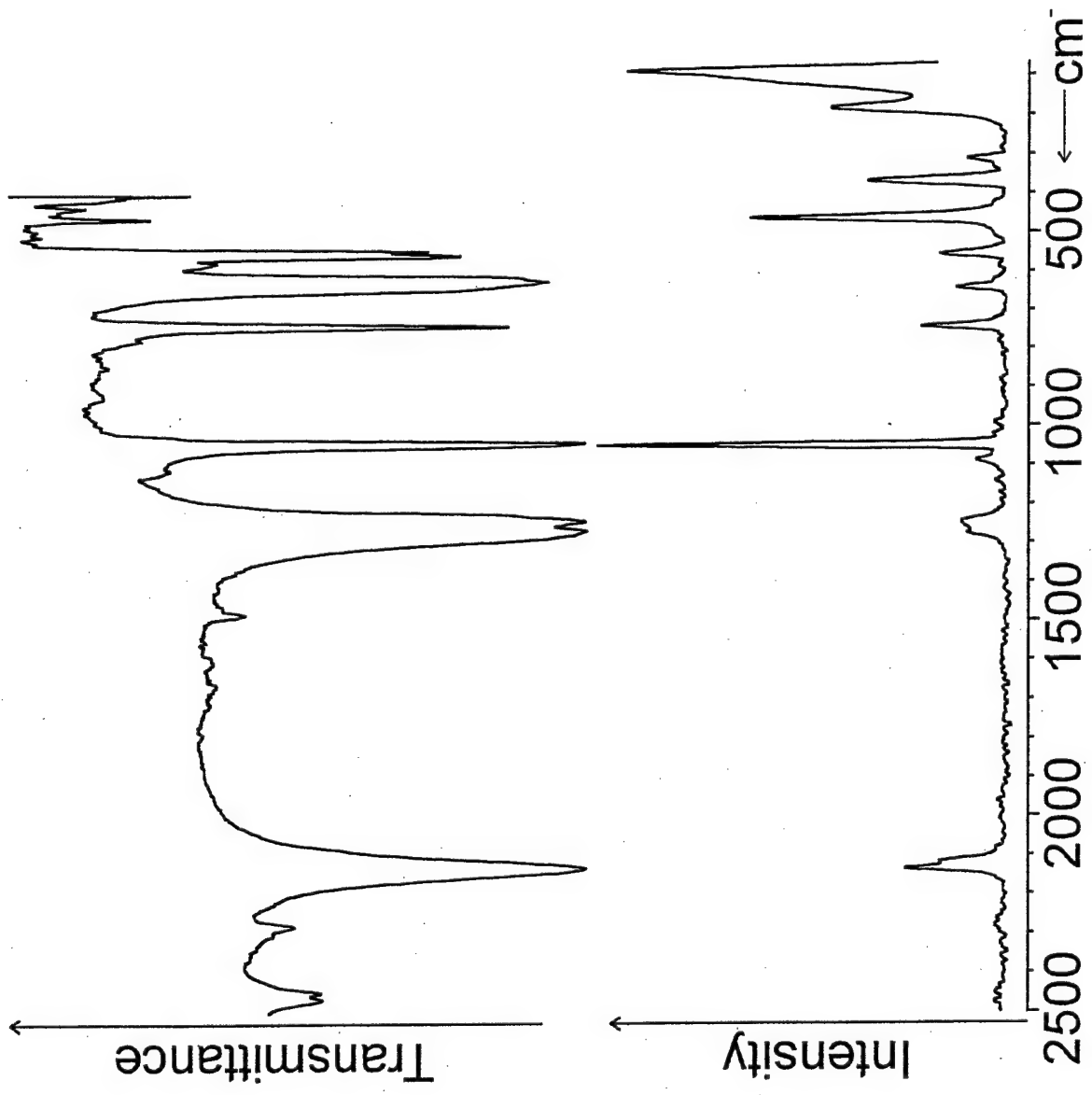




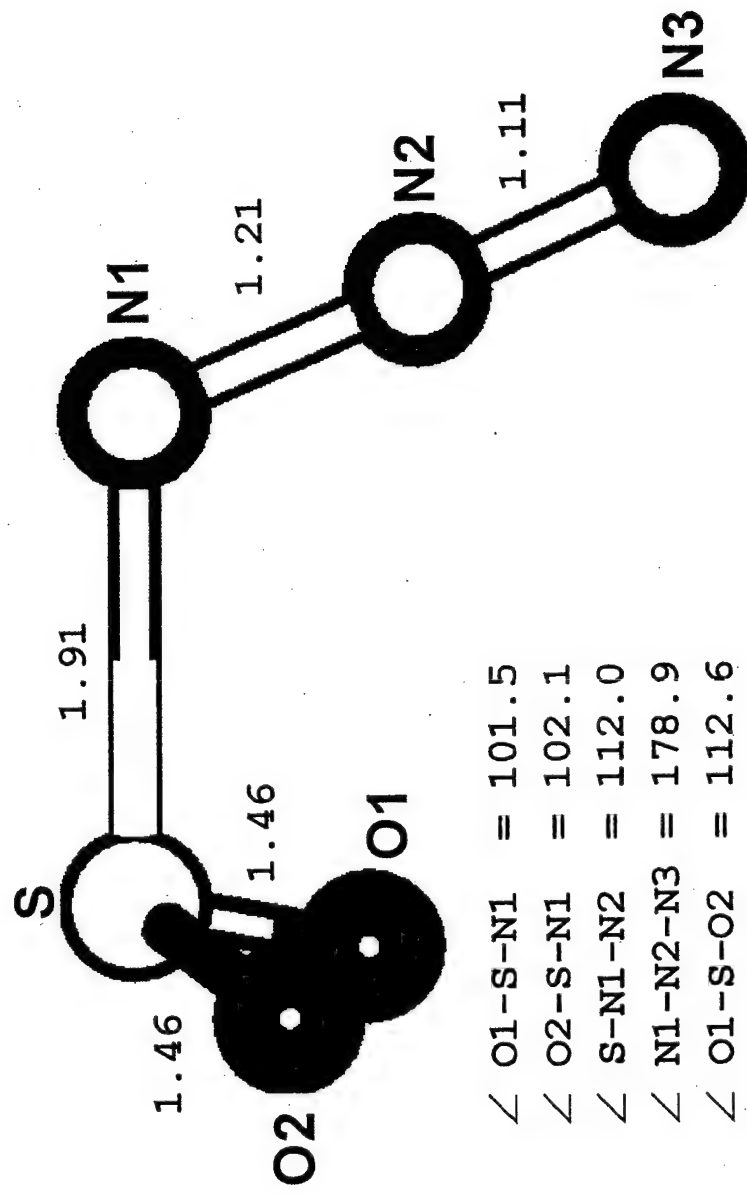






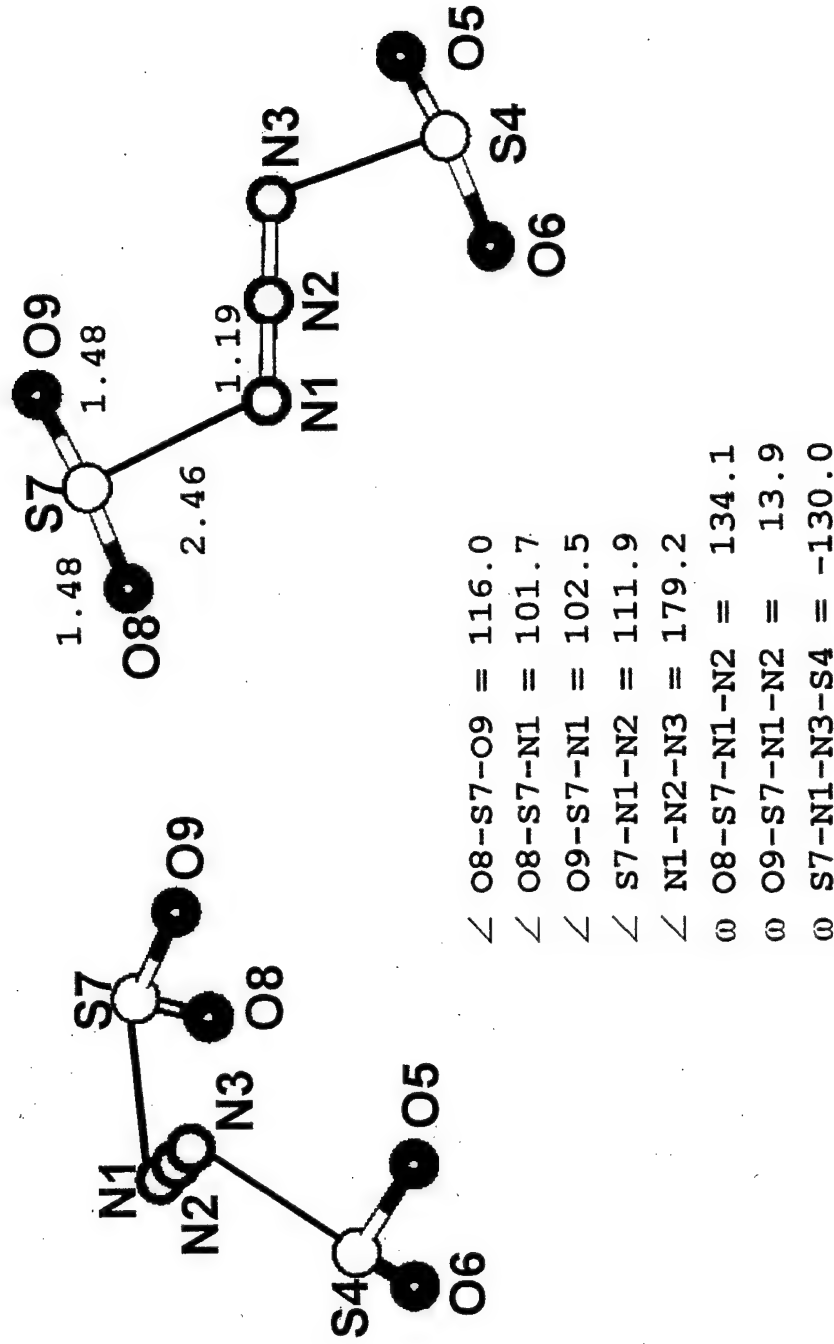


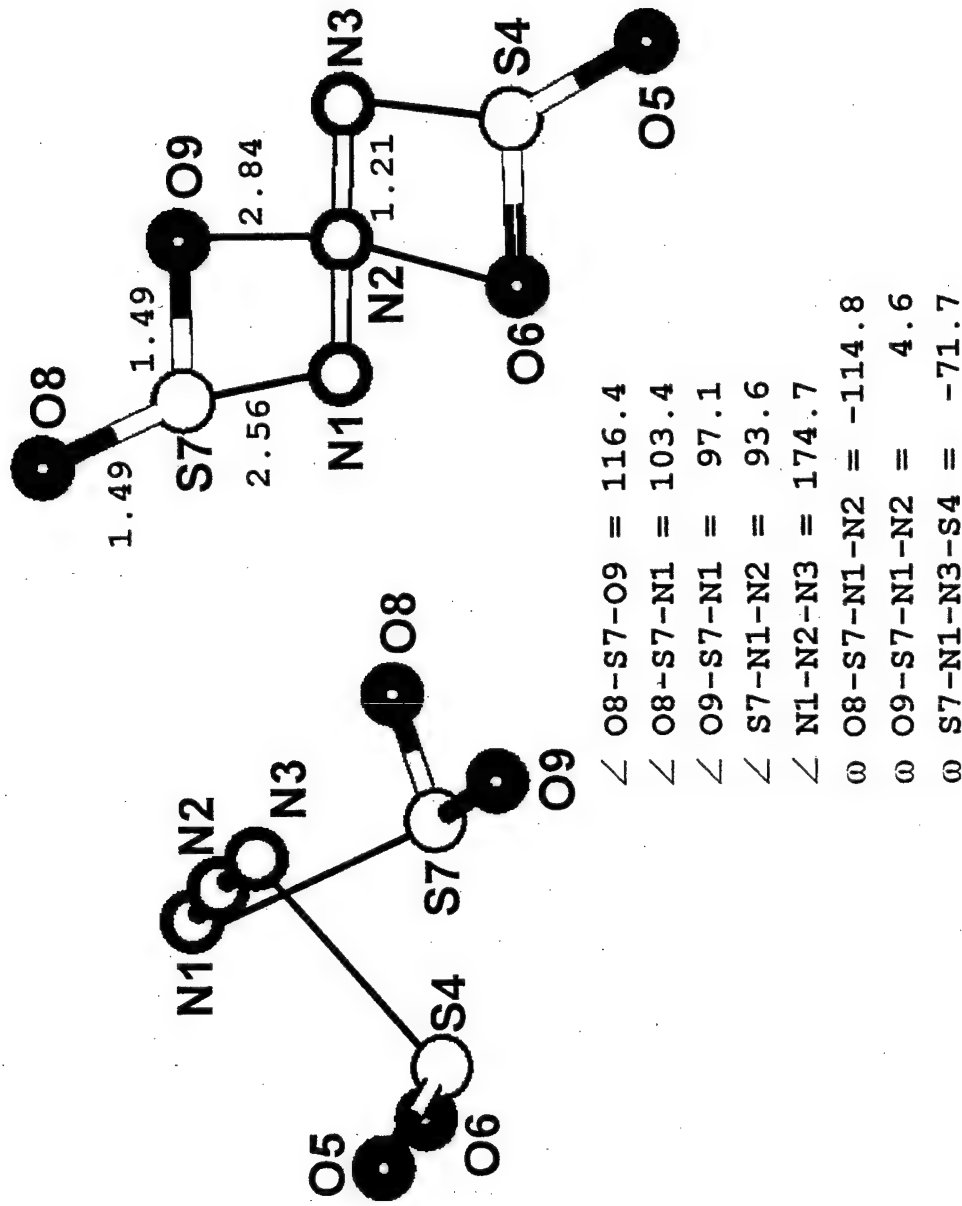
RHF/6-31+G (d)

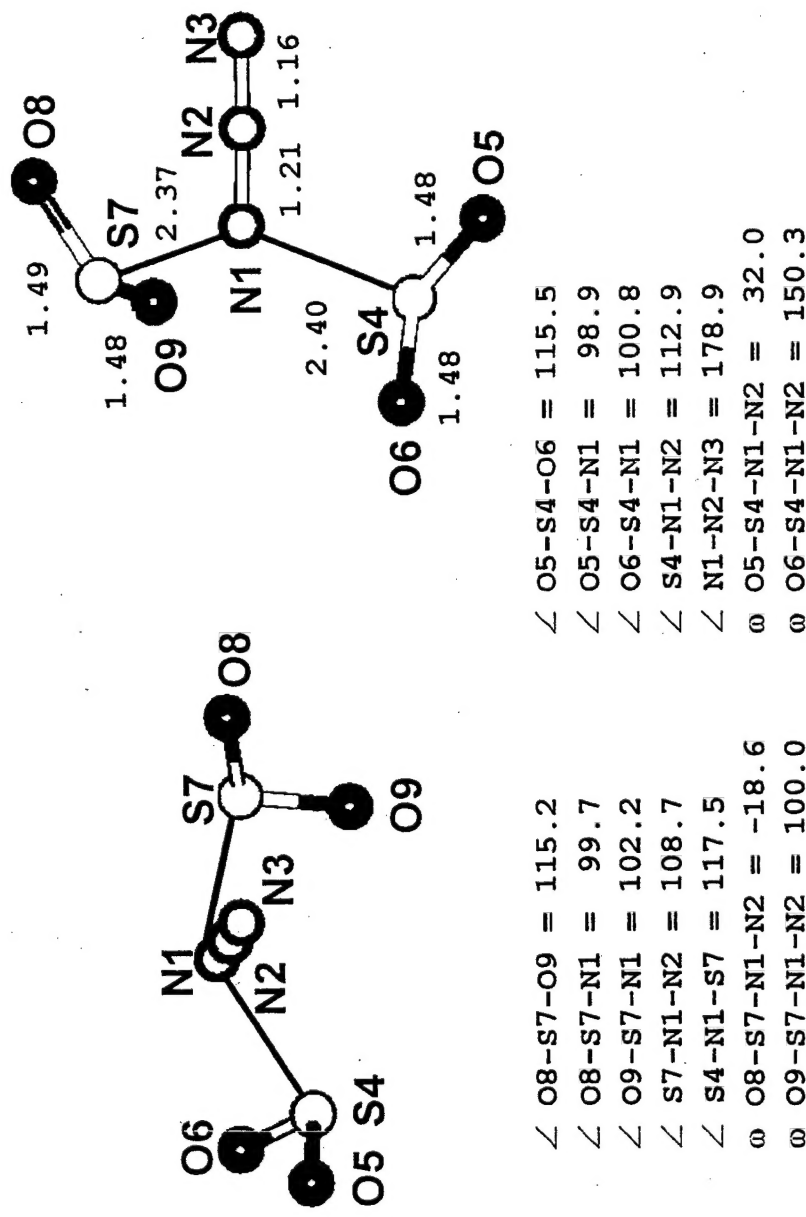


$\angle O1-S-N1 = 101.5$
 $\angle O2-S-N1 = 102.1$
 $\angle S-N1-N2 = 112.0$
 $\angle N1-N2-N3 = 178.9$
 $\angle O1-S-O2 = 112.6$

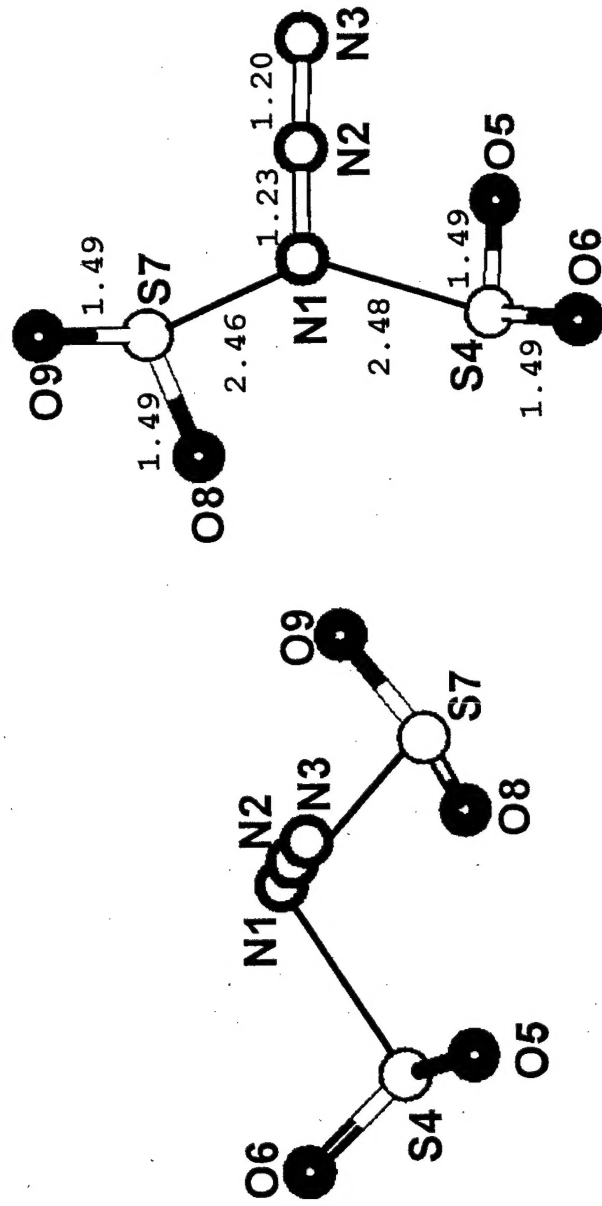
B3LYP/6-31+G(d)







MP2/6-31+G(d)



\angle O8-S7-O9 = 116.3
 \angle O8-S7-N1 = 100.3
 \angle O9-S7-N1 = 101.2
 \angle S7-N1-N2 = 102.3
 \angle S4-N1-S7 = 100.7
 ϖ O8-S7-N1-N2 = 147.3
 ϖ O9-S7-N1-N2 = -93.1
 \angle O5-S4-N1-N2 = 116.3
 \angle O5-S4-N1 = 97.3
 \angle O6-S4-N1 = 102.1
 \angle S4-N1-N2 = 99.3
 \angle N1-N2-N3 = 177.5
 ϖ O5-S4-N1-N2 = -19.5
 ϖ O6-S4-N1-N2 = 99.5

Karl O. Christe,^{*†} Jerry A. Boatz,[†] Michael Gerken,[†] Ralf Haiges,^{*} Stefan Schneider,[‡] Thorsten Schroer,^{*} Fook S. Tham,[†] Ashwani Vij,[†] Vandana Vij,[†] Ross I. Wagner,^{*} and William W. Wilson[†]

Inorg. Chem. 2002, 41, ...
Synthesis and Characterization of the
 SO_2N_3^- , $(\text{SO}_2)_2\text{N}_3^-$, and SO_3N_3^- Anions

The azide anion forms colorless 1:1 and yellow 1:2 adducts with SO_2 . These adducts were characterized by vibrational and NMR spectroscopy and theoretical calculations. The crystal structure of $\text{CsSO}_2\text{N}_3\text{CsSO}_3\text{N}_3^-$ was determined and shows that the N-S bond in SO_2N_3^- is 23 Å longer than that in SO_3N_3^- . Both N-S bonds are unusually long, resulting in facile SO_2 ligand exchange and low barriers toward rotational disorder.

